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(54) Title: MAMMALIAN EDG-5 RECEPTOR HOMOLOGS

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(57) Abstract

The present invention is directed to nucleic seld sequences and amino acid sequences for mammalian EDG-5 receptor homologs, and particularly for human EDG-5 receptor homologs. The invention also provides methods for determining agonists and antagonists for EDG-5 receptors in addition to assays, expression vectors, host cells and methods for treating disorders associated with abernant expression or activity of EDG-5. SIP and SPC are agonists for EDG-5 receptors.

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MAMMALIAN EDG-5 RECEPTOR HOMOLOGS

FIELD OF THE INVENTION

S mammalian, including human, EDG-5 receptor homologs. present invention describes a nucleic acid sequence and an amino acid sequence for novel The present invention is in the field of molecular biology; more particularly, the

BACKGROUND OF THE INVENTION

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ᅜ Biol. 135: 1071-1083; An et al. (1997) Biochem. Biophys. Res. Comm. 213: 619-622). Chem. 265:9308-13; US 5,585,476). Recently, however, lysophospatidic acid has been demonstrated to be the endogenous ligand for the edg-2 receptor (Hecht et al. (1996) J. Cell their endogenous ligands are not known (for example see HIa T and Maciag T (1990) J Biol The family of edg receptors are commonly grouped with orphan receptors because

20 22 ö receptor. Specifically: the TM-VII is generally a highly conserved portion of the T7G extracellular N-terminal segment or one or more of the three extracellular loops participate in segments (TMS) are designated by roman numerals I-VII and account for structural and plasma membrane and form a bundle of antiparallel α helices. These transmembrane (T7Gs). T7Gs are so named because of their seven hydrophobic domains, which span the binding and in subsequent induction of conformational change in intracellular portions of the pocket; however, when the binding site must accommodate more bulky molecules, the shows greatly reduced conservation receptors, and is often critically involved in ligand binding and receptor activation. the functional features of the receptor. In most cases, the bundle of helices forms a binding usually hydrophilic and highly antigenic relative to the receptor polypeptide as a whole and those that transduce intracellular signals upon receptor activations; the carboxy-terminal is intracellular carboxy-terminal is involved in interactions with intracellular proteins, including The edg family of receptors is seven transmembrane G protein coupled receptors

of protein kinases, alteration in the expression of specific genes. messengers such as cyclic AMP (cAMP), phospholipase C, inositol triphosphate, activation which mediates further intracellular signaling activities generally the production of second Once the receptor is activated, it interacts with an intracellular G-protein complex

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differentially modulate distinct intracellular pathways that are controlled from T7G receptors physiological or pharmaceutical molecules, which trigger, prolong or inhibit its activity or or intervene in such processes, and the receptor can be used in screening assays to identify disease processes. Identification of a novel T7G receptor provides the opportunity to diagnose T7G receptors are expressed and activated during numerous developmental and

SUMMARY OF THE INVENTION

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EDG-5 (HEDG-5-5). Herein, the nucleotide sequence encoding MEDG-5 and HEDG-5 is mammalian receptor homologs EDG-5, including murine EDG-5 (MEDG-5) and human designated medg-5 and hedg-5, respectively. The invention provides isolated and unique nucleotide sequences which encode novel

nucleotide sequences which hybridize under stringent conditions to edg-5, particularly, hedg the complement of edg-5 mRNA, particularly hedg-5. In addition, the invention features The present invention also relates to the isolated and unique nucleotide sequences of

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20 comprising such edg-5 nucleotide sequences In addition, the present invention relates to expression vectors and host cells

antibodies for EDG-5, for example fragments in the TM-VII and carboxy-terminal domain. More particularly, the present invention provides fragments which are useful as

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of mammalian EDG-5, and more particularly, to the use of HEDG-5, or its variants, in the diagnosis or treatment of diseased cells and/or tissues associated with aberrant expression of Furthermore, the invention relates to the use of nucleic acid and amino acid sequences

or expression vectors containing edg-5/hedg-5; host cells or organisms transformed with particularly, the HEDG-5 receptor, and include: the antisense DNA of edg-5/hedg-5; cloning expression vectors containing edg-5/hedg-5; chromosomal localization of hedg-5; expression Additional aspects of the invention are directed to the EDG-5 receptor, but more

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and tissue distribution of edg-5/hedg-5; a method for the production and recovery of purified EDG-5/HEDG-5 from host cells; purified protein, EDG-5/HEDG-5, which can be used to identify inhibitors for the downregulation of signal transduction involving EDG-5/HEDG-5; and methods of screening for ligands of edg-5/hedg-5 using transformed cells.

Particularly there is provided an isolated nucleotide sequence selected from the group consisting of:

- (a) the nucleotide sequence comprising nucleotides 36-10974 of SEQ. ID NO: 13
- (Figure 3A) (b) the nucleotide sequence of Figure 3B;

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- (c) the nucleotide sequence of Figure 3C;(d) the nucleotide sequence comprising at least about 70% sequence identity to (a),
- (b) or (c), more preferably at least about 80-85% sequence identity, and even more preferably at least about 90% sequence identity, and most preferably at least about 95% sequence identity, and which nucleotide sequence hybridizes under stringent conditions to the
- nucleotide sequence of (a), (b) or (c), respectively; or portions thereof, and
- (e) the nucleotide sequence which encodes the amino acid sequence of Figure 4A (SEQ ID NO. 14), 4B or 4C. There is also provided: expression vectors; host cells; purified amino acid sequences; complementary nucleic acid sequences; biologically active fragments; and hybridization probes, for such nucleotide sequences and their encoded amino acid
- 20 sequences.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A shows a partial DNA sequence of clone 501 which is a murine edg-5 clone (SEQ ID NO: 3).

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Figure 1B shows the full length DNA sequence of the a subclone of a mhedg-5 pBluescript subclone and the predicted amino acid sequence thereof.

30 Figure 2 shows the amino acid sequence encoded by the DNA sequence of Figure 1A (SEQ ID NO: 15)

Figure 3A shows a nucleotide sequence of hedg-5 cDNA inserted into pcDNA3, nucleotides 36-1097 of which encode the full length HEDG-5. (pC3-hEdg5-3)

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Figure 3B shows a nucleotide sequence of hedg-5 cDNA of clone pC3-hEdg5#3.4, which encodes the full length HEDG-5.

Figure 3C shows a nucleotide sequence of hedg-5 cDNA of clone pc3-hEdg5#28, which encodes the full length HEDG-5.

Figure 4A shows an alignment of the genomic DNA of Figure 3A (which corresponds to the cDNA of the pC3-hEdg5-3 from nt 251-1523 and the genomic DNA flanking from nt 1-250) with the predicted amino acid sequence.

Figure 4B shows the predicted amino acid sequence of hedg-5 cDNA of Figure 3B.

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Figure 4C shows the predicted amino acid sequence of hedg-5 cDNA of Figure 3C.

Figure 5A shows the alignment of the predicted amino acid sequences of HEDG5 translation products of clones pC3-hedg5-3, pC3-hedg5#4, and pC3-hedg5#28 as set out in Figures 4A, 4B and 4C, respectively.

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Figure 5B shows the alignment of the amino acid sequence of murine edg-5 with the animo acid sequence of human edg-5 from the pC3-hEdg5#3.4 clone.

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Figure 6 shows the functional response of the pC3-hedg5#4, pC3-hedg5-3 and pC3-hedg5#28 clones to anandamide and to LPA by activation of NF-kB production.

25 Figure 7 shows the SRE response and AP-1 response of pC3-hedg5#28 when treated with 10 μM LPA.

DETAILED DESCRIPTION OF THE INVENTION

The invention relates in one respect to polynucleotides, in their isolated form, that code for mammalian, including murine and human, EDG-5 receptors. The EDG receptors are characterized by structural features common to the G-protein coupled receptor class,

of a G-protein coupled receptor such as an EDG-5 receptor can be measured using any of a lysophospholipid selectively. When expressed functionally in a host cell, i.e., in operable including seven transmembrane regions, and by the functional properties of binding variety of appropriate functional assays described hereinbelow. of responding to lysophospholipid binding by signal transduction. In this regard, the activity linkage with a responsive second messenger system the EDG-5 receptors are capable further

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mammalian receptor, EDG-5, are characterised by structural features common to the Gactive fragments thereof and the lower case edg-5 to the nucleotide sequence thereof. The mammalian EDG-5 receptor homolog in either naturally occurring or synthetic form and protein coupled receptors, including the seven transmembrane regions, and by the sequence identity to each other of greater than about 56%, more preferably greater than about 70% identity, and most preferably greater than about 80% identity. As used herein and designated by the upper case abbreviation, EDG-5, refers to

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and the nucleotide sequence as medg-5. the nucleotide sequence as hedg-5 and the murine EDG-5 receptor is designated as MEDG-5 Furthermore, as used herein, the human EDG-5 receptor is designated as HEDG-5 and

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25 nucleotide level. In the studies detailed herein the hedg-5 sequence was used, however, these comparison with known sequences demonstrated that this mouse clone represented a gene domains (TM-2) and TM-7 of the G protein-coupled receptor (GPCR) superfamily. Sequence neuronal cell line using degenerate primers based on conserved regions of transmembrane sequence disclosed herein studies and the applications detailed herein could be undertaken using the novel mouse edg-5 related to, but not identical to edg-2, an orphan GPCR. Sequence identity was 49% at the The novel murine hedg-5 sequence was isolated following PCR from a murine

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in the art. For example, by screening an arrayed mouse library (Genome Systems Inc.) using stringency, with the final wash done for 30 min at 65 C in 1X SSC. Genomic DNA inserts condon priming method and then hybridized to the PAC filters and washed at high the full-length human edg-5 cDNA. The hedg-5 sequence is first radiolabelled using the A full-length mouse sequence is obtained using methods well known to those of skill

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comparable cloning vector, using at least 3 different restriction digests of which 1 should human edg-5 sequences. The position of the single intron seen in the human edg-5 gene picked, grown, mapped by restriction digest and Southern blotting to identify the size of the be screened with the same cDNA probe under the same stringency conditions. Positives are have a 4 bp recognition site. Each digest yields a different subclone library, which in turn can from the clones with the strongest signals can be shotgun subcloned into pBluescript or a confidence to obtain the complete coding sequence of the mouse edg-5 gene without should be conserved in the mouse gene. Thus, primers can be designed with a high degree of hybridizing insert, then sequenced using primers based on either the vector sequence, or on

5 a Laboratory Manual (Cold Spring Harbor Press, 1989). more detailed description of this approach can be found in Manialis et al. Molecular Cloning, can be designed to amplify the cDNA directly from various tissue and/or cell line sources. A including intron sequences. Once the coding region has been determined, new PCR primers

reference for the purpose of describing the methodologies, cell lines and vectors, among other entitled to antedate such disclosure, for example, by virtue of prior invention things. However, nothing herein is to be construed as an admission that the invention is not All publications and patent applications mentioned herein are incorporated by

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20 Definitions

pertains. terms used in the application. Any terms not specifically defined should be given the meaning commonly understood by one of oridinary skill in the art to which the invention The following definitions are used herein for the purpose of describing particular

form of RNA, or in the form of DNA including cDNA, genomic DNA and synthetic DNA. polynucleotide is considered "isolated" when it has been selected, and hence removed from proteins. In the context of polynucleotide libraries, for instance, a EDG-5 receptor-encoding association with other polynucleotides within the library. Such polynucleotides may be in the As used herein "isolated" means separated from polynucleotides that encode other

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environment, and are isolated or separated, and are at least 60% free, preferably 75 % free, As used herein "purified" refers to sequences that are removed from their natural

and most preferably 90% freee from other components with which they are naturally ussociated.

An "oligonucleotide" is a stretch of nucleotide residues which has a sufficient number of bases to be used as an oligomer, amplimer or prohe in a polymerase chain reaction (PCR). Oligonucleotides are prepared from genomic or cDNA sequence and are used to amplify, reveal or confirm the presence of a similar DNA or RNA in a particular cell or tissue. Oligonucleotides or oligomers comprise portions of a DNA sequence having at least about 10 nucleotides and as many as about 35 nucleotides, preferably about 25 nucleotides.

"Probes" may be derived from naturally occurring or recombinant single - or double - stranded nucleic acids or be chemically synthesized. They are useful in detecting the presence of identical or similar sequences.

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A "portion" or "fragment" of a polynucleotide or nucleic acid comprises all or any part of the nucleotide sequence having fewer nucleotides than about 6 kb, preferably fewer than about 1 kb. A portion or fragment can be used as a probe. Such probes may be labeled with reporter molecules using nick translation, Klenow fill-in reaction, PCR or other methods well known in the art. To optimize reaction conditions and to eliminate false positives, well known in the art. To optimize reaction conditions and to eliminate false positives, whether DNA or RNA encoding HEDG-5 is present in a cell type, tissue, or organ.

"Reporter" molecules are those radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents which associate with, establish the presence of, and may allow quantification of a particular nucleotide or amino acid sequence.

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"Recombinant nucleotide variants" encoding HEDG-5 may be synthesized by making use of the "redundancy" in the genetic code. Various codon substitutions, such as the silent changes which produce specific restriction sites or codon usage-specific mutations, may be introduced to optimize cloning into a plasmid or viral vector or expression in a particular prokaryotic or eukaryotic host system, respectively.

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"Chimeric" molecules may be constructed by introducing all or part of the nucleotide sequence of this invention into a vector containing additional nucleic acid sequence which might be expected to change any one (or more than one) of the following HEDG-5 characteristics: cellular location, distribution, ligand-binding affinities, interchain affinities, degradation/turnover rate, signaling, etc.

"Biologically Active or Active" refers to those forms, fragments, or domains of any HEDG-5 polypeptide which retain at least some of the biological and/or antigenic activities of any naturally occurring HEDG-5.

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"Naturally occurring HEDG-5" refers to a polypeptide produced by cells which have not been genetically engineered and specifically contemplates various polypeptides arising from post-translational modifications of the polypeptide including but not limited to acctylation, carboxylation, glycosylation, phosphorylation, lipidation and acylation.

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"Derivative" refers to those amino acid and nucleotide sequences which have been chemically modified. Such techniques for amino acid deravitives include: ubiquitination; labeling (see above); pegylation (derivatization with polyethylene glycol); and chemical insertion or substitution of amino acids such as ornithine which do not normally occur in human proteins. A nucleotide sequence derivative would encode the amino acid which retains its essential biological characteristics of the natural molecule.

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"Recombinant polypeptide variant" refers to any polypeptide which differs from naturally occurring HEDG-5 by amino acid insertions, deletions and/or substitutions, created using recombinant DNA techniques. Guidance in determining which amino acid residues may be replaced, added or deleted without abolishing activities of interest may be found by comparing the sequence of HEDG-5 with that of related polypeptides and minimizing the number of amino acid sequence changes made in highly conserved regions.

Amino acid "substitutions" are conservative in nature when they result from replacing one amino acid with another having similar structural and/or chemical properties, such as the replacement of a leucine with an isoleucine or valine, an aspartate with a glutamate, or a threonine with a serine.

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"Insertions" or "deletions" are typically in the range of about 1 to 5 amino acids. The variation allowed may be experimentally determined by producing the peptide synthetically or by systematically making insertions, deletions, or substitutions of nucleotides in the hedg-5 sequence using recombinant DNA techniques.

A "signal or leader sequence" can be used, when desired, to direct the polypeptide through a membrane of a cell. Such a sequence may be naturally present on the polypeptides of the present invention or provided from heterologous sources by recombinant DNA

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An "oligopeptide" is a short stretch of amino acid residues and may be expressed from an oligonucleotide. It may be functionally equivalent to and the same length as (or considerably shorter than) a "fragment", "portion", or "segment" of a polypeptide. Such sequences comprise a stretch of amino acid residues of at least about 5 amino acids and often about 17 or more amino acids, typically at least about 9 to 13 amino acids, and of sufficient length to display biological and/or antigenic activity.

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"Inhibitor" is any substance which retards or prevents a biochemical, cellular or physiological reaction or response. Common inhibitors include but are not limited to antisense molecules, antibodies, and antagonists.

"Standard" is a quantitative or qualitative measurement for comparison. It is based on a statistically appropriate number of normal samples and is created to use as a basis of comparison when performing diagnostic assays, running clinical trials, or following patient treatment profiles.

"Stringent conditions" is used herein to mean conditions that allow for hybridization of substantially related nucleic acid sequences. Such hybridization conditions are described by Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, 1989. Generally, stringency occurs within a range from about 5 °C below the melting temperature of the probe to about 20 °C – 25 °C below the melting temperature. As understood by ordinary skilled persons in the art, the stringency conditions may be altered in

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order to identify or detect identical or related nucleotide sequences. Factors such as the length and nature (DNA, RNA, base composition) of the sequence, nature of the target (DNA, RNA, base composition, presence in solution or immobilization, etc.) and the concentration of the salts and other components (e.g. the presence or absence of formamide, dextran sulfate and/or polyethylene glycol) are considered and the hybridization solution may be varied to generate conditions of either low or high stringency.

"Animal" as used herein may be defined to include human, domestic (cats dogs, etc.), agricultural (cows, horses, sheep, etc.) or test species (mouse, rat, rabbit, etc.).

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"Nucleotide sequences" as used herein are oligonucleotides, polynucleotides, and fragments or portions thereof, and are DNA or RNA of genomic or synthetic origin which may be single or double stranded, and represent the sense or complement or antisense strands

"Sequence identity" is known in the art, and is a relationship between two or more polypeptide sequences or two or more polynucleotide sequences, as determined y comparing the sequences, particularly, as determined by the match between strings of such sequences. Sequence identity can be readily calculated by known methods (Computational Molecular Biology, Lesk, A.M., ed., Oxford University Press, New York, 1988; Biocomputing:

Informatics and Genome Projects, Smith, D.W., ed., Academic Press, New York, 1993;

Computer Analysis of Sequence Data, Part I, Griffin, A.M., and Griffin, H.G., eds., Humana Press, New Jersey, 1994; Sequence Analysis in Molecular Biology, von Heinje, G., Academic Press, 1987; and Sequence Analysis Primer, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991). While there exist a number of methods to measure identity between two sequences, the term is well known to skilled artisans (see, for example, Sequence Analysis in Molecular Biology; Sequence Analysis Primer; and Carillo, H., and Lipman, D.,

DNASIS (Hitachi Software Engineering Co., San Bruno, CA). Computer programs for determining identity are publicly available. Preferred computer program methods to determine identity between two sequences include, but are not limited to, GCG program

SIAM J. Applied Math., 48: 1073 (1988)). Methods commonly employed to determine identity between sequences include, but are not limited to those disclosed in Carillo, H., and

Lipman, D., SIAM J. Applied Math., 48: 1073 (1988) or, preferably, in Needleman and Wunsch, J. Mol. Biol., 48: 443-445, 1970, wherein the parameters are as set in version 2 of

package (Devereux, J., et al., Nucleic Acids Research 12(1): 387 (1984)), BLASTP, BLASTN, and FASTA (Atschul, S.F. et al., J. Molec. Biol. 215: 403-410 (1990)). The BLASTX program is publicly available from NCBI (blast@ncbi.nlm.nih.gox) and other sources (BLAST Manual, Altschul, S., et al., NCBI NLM NIH Bethesda, MD 20894; Altschul, S., et al., J. Mol. Bio. 215: 403-410 (1990)). Computational Molecular Biology, Lesk, A.M., ed. Unless specified otherwise in the claims, the percent identity for the purpose of interpreting the claims shall be calculated by the Needleman and Wucnsch algorithm with the parameters set in version 2 of DNASIS.

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The present invention provides a nucleotide sequence uniquely identifying novel mammalian, including murine (MEDG-5) and human (HEDG-5), seven transmembrane receptor (T7G) or EDG-5.

Based on the homology of HEDG-5 to human edg-2 (see table 2 below) it is likely that HEDG-5 binds a ligand of the same chemical class. Edg-2 specifically binds lysophosphatidic acid (LPA) which is a phospholipid. It was determined herein that HEDG-5 also recognizes LPA as a functional agonist.

20 25 30 including mitogenisis (Xu et al. (1995) J. Cell. Physiol., 163: 441-450) and apoptosis, cell (1992) J. Biol. Chem., 267: 21360-21367). Further, considerable circumstantial evidence and cell migration (Imamura et al. (1993) Biochem. Biophys. Res. Comm., 193: 497-503). It factor-like effects on cell proliferation (Moolenar (1996) J. Biol. Chem, 270: 12949-12952) adhesion and regulation of gene expression. Specifically, for example, LPA elicits growth rheumatoid arthritis (Natiarajan et al. (1995) J. Lipid Res., 36(9): 2005-2016), psoriasis and has also been suggested that LPA plays a role in wound healing and regeneration (Tigyi et al. of HEDG-5 is likely to be associated with a chronic or acute disease states. Further, inflammatory bowel disease. Thus, ligands for HEDG-5, including LPA, are likely to be respiratory distress, neurodegeneration (Jalink et al. (1993) Cell Growth Differ., 4: 247-255) inflammatory component (Fourcade et al. (1995), Cell, 80(6): 919-927, including adult (Imamura et al., (1993) Biochem. Biophys. Res. Comm., 193: 497-503); diseases having an indicates that phospholipids may be involved in various disease states including cancer biologically important regulators of cell activity, and therefore aberrant expression or activity Phospholipids have been demonstrated to be important regulators of cell activity,

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modulators of HEDG-5 activity are likely to be useful in treatment or prevention of such disease states.

5 2 example, preparation of fresh stock solutions (e.g., 10 mM) by dissolving the phospholipid ir of compounds.. Therefore, in one embodiment, preferably phospholipid molecules should be metabolic stability, for example, by changing ester bond at Sn-1 to an ether on by blocking activating factor (lyso-PAF) and phosphatidic acid. These ligands can be altered to improve (LPS), lysophosphatidylinositol (LPI), lysophosphatidylcholine (LPC), lyso-platelet should be screened, such as lysophosphatidylethanolamine (LPE), lysophosphatidylserine screened to identify HEDG-5 ligands. Even more preferably, lysoglycerophospholipids phosphate. LPA and related phospholipids have limited solubility in aqueous solution and benefits may be derived from shortening the fatty acid chain or altering the positioning of the the free hydroxyl group with methoxy or acetyl ester. Additional medicinal chemistry calcium-free PBS and faity-acid free BSA. Other related phospholipids can be prepared, for have a tendency to be sticky. These problems may be alleviated in a number of ways. For example, in 100% ethanol or DMS0 HEDG-5 ligands, other than LPA, are likely to be found among the phospholipid class

A diagnostic test for aberrant expression of HEDG-5 can accelerate diagnosis and proper treatment of abnormal conditions of for example, the heart, kidney, lung and testis. Specific examples of conditions in which aberrant expression of HEDG-5 may play a role include adult respiratory distress, asthma, rheumatoid arthritis, cardiac ischemia, acute pancreatitis, septic shock, psoriasis, acute cyclosporine nephrotoxicity and early diabetic glomerulopathy, as well as lung damage following exposure to cigarette smoke, asbestos or silica.

The nucleotide sequences encoding EDG-5 (or their complement) have numerous applications in techniques known to those skilled in the art of molecular biology. These techniques include use as hybridization probes, use in the construction of oligomers for PCR.

30 use for chromosome and gene mapping, use in the recombinant production of EDG-5, and use in generation of antisense DNA or RNA, their chemical analogs and the like. Uses of nucleotides encoding EDG-5 disclosed herein are exemplary of known techniques and are not intended to limit their use in any technique known to a person of ordinary skill in the art.

Furthermore, the nucleotide sequences disclosed herein may be used in molecular biology techniques that have not yet been developed, provided the new techniques rely on properties of nucleotide sequences that are currently known, e.g., the triplet genetic code, specific base pair interactions, etc.

It will be appreciated by those skilled in the art that as a result of the degeneracy of the genetic code, a multitude of EDG-5-encoding nucleotide sequences may be produced. Some of these will only bear minimal homology to the nucleotide sequence of the known and naturally occurring EDG-5. The invention has specifically contemplated each and every possible variation of nucleotide sequence that could be made by selecting combinations based on possible codon choices. These combinations are made in accordance with the standard triplet genetic code as applied to the nucleotide sequence of naturally occurring edg-5, and all such variations are to be considered as being specifically disclosed.

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Although the nucleotide sequences which encode EDG-5, its derivatives or its variants are preferably capable of hybridizing to the nucleotide sequence of the naturally occurring edg-5 under stringent conditions, it may be advantageous to produce nucleotide sequences encoding EDG-5 or its derivatives possessing a substantially different codon usage. Codons can be selected to increase the rate at which expression of the peptide occurs in a particular particular codons are utilized by the host. Other reasons for substantially altering the nucleotide sequence encoding EDG-5 and/or its derivatives without altering the encoded as sequence include the production of RNA transcripts having more desirable properties, such as a greater half-life, than transcripts produced from the naturally occurring sequence.

Human genes often show considerable actual polymorphism; that is, variation in nucleotide sequence among a fraction of the entire human population. In many cases this polymorphism can result in one or more amino acid substitutions. While some of these substitutions show no demonstrable change in function of the protein, others may produce varying degrees of functional effects. In fact, many natural or artificially produced mutations can lead to expressible HEDG proteins. Each of these variants, whether naturally or artificially produced, is considered to be equivalent and specifically incorporated into the

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Nucleotide sequences encoding EDG-5 may be joined to a variety of other nucleotide sequences by means of well established recombinant DNA techniques (Sambrook J et al (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold

- 5 (1989) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor NY; or Ausubel FM et al (1989) Current Protocols in Molecular Biology, John Wiley & Sons, New York City). Useful nucleotide sequences for joining to edg-5 include an assortment of cloning vectors such as plasmids, cosmids, lambda phage derivatives, phagemids, and the like. Vectors of interest include expression vectors, replication vectors,
- probe generation vectors, sequencing vectors, etc. In general, vectors of interest may contain an origin of replication functional in at least one organism, convenient restriction endonuclease sensitive sites, and selectable markers for one or more host cell systems.

Another aspect of the subject invention is to provide for edg-5-specific hybridization probes capable of hybridizing with naturally occurring nucleotide sequences encoding EDG-5. Such probes may also be used for the detection of similar T7G encoding sequences and should preferably contain at least 56% nucleotide identity, more preferably at least 70% identity, to edg-5 sequence. The hybridization probes of the subject invention may be derived from the nucleotide sequence presented as SEQ. ID NO:12 or from genomic sequences

Hybridization probes may be labeled by a variety of reporter molecules using techniques well known in the art. Preferably, the hybridization probes incorporate at least 15 nucleotides, and preferably at least 25 nucleotides, of the edg-5 receptor, more particualry of the medg-5 or the hedg-5 receptor. Suitable hybridization probes would include: consensus fragments, i.e. those regions of the mouse and human edg-5 receptor that are identical (See Figure 5B); the extracellular edg-5 binding domain, the stipulated transmembrane regions and the C-terminal

It will be recognized that many deletional or mutational analogs of nucleic acid sequences for EDG-5 will be effective hybridization probes for EDG-5 nucleic acid.

portion of the receptor

30 Accordingly, the invention relates to nucleic acid sequences that hybridize with such EDG-5 encoding nucleic acid sequences under stringent conditions.

"Stringent conditions" refers to conditions that allow for the hybridization of substantially related nucleic acid sequences. For instance, such conditions will generally allow hybridization of sequence with at least about 70% identity, preferably with at least 80-85% sequence identity, more preferably with at least about 90% sequence identity, and even more preferably with at least about 95% sequence identity. Such hybridization conditions are described by Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, 1989. Hybridization conditions and probes can be adjusted in well-characterized ways to achieve selective hybridization of human-derived probes. Nucleic acid molecules that will hybridize to EDG-5 encoding nucleic acid under stringent conditions can be identified functionally, using methods outlined above, or by using for example the

hybridization rules reviewed in Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, 1989. Without limitation, examples of the uses for hybridization probes include: histochemical uses such as identifying tissues that express EDG-5; measuring mRNA levels, for instance to identify a sample's tissue type or to identify cells that express abnormal levels of EDG-5; and detecting polymorphisms in the EDG-5.

RNA hybridization procedures are described in Maniatis et al. Molecular Cloning, a Laboratory Manual (Cold Spring Harbor Press, 1989). PCR as described US Patent No's. 4,803,195; 4,800,195; and 4,965,188 provides additional uses for oligonucleotides based upon the nucleotide sequence which encodes the edg-5 sequences of the invention. Such

probes used in PCR may be of recombinant origin, chemically synthesized, or a mixture of both. Oligomers may comprise discrete nucleotide sequences employed under optimized conditions for identification of edg-5 in specific tissues or diagnostic use. The same two oligomers, a nested set of oligomers, or even a degenerate pool of oligomers may be employed under less stringent conditions for identification of closely related DNA's or RNA's. Rules for designing PCR primers are now established, as reviewed by PCR

Protocols, Cold Spring Harbor Press, 1991. Degenerate primers, i.e., preparations of primers that are heterogeneous at given sequence locations, can be designed to amplify nucleic acid sequences that are highly homologous to, but not identical to edg-5. Strategies are now available that allow for only one of the primers to be required to specifically hybridize with a

30 known sequence. See, Froman et al., Proc. Natl. Acad. Sci. USA 85: 8998, 1988 and Loh et al., Science 243: 217, 1989. For example, appropriate nucleic acid primers can be ligated to the nucleic acid sought to be amplified to provide the hybridization partner for one of the primers. In this way, only one of the primers need be based on the sequence of the nucleic

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acid sought to be amplified. PCR methods of amplifying nucleic acid will utilize at least two primers. One of these primers will be capable of hybridizing to a first strand of the nucleic acid to be amplified and of priming enzyme-driven nucleic acid synthesis in a first direction.

The other will be capable of hybridizing the reciprocal sequence of the first strand (if the sequence to be amplified is single stranded, this sequence will initially be hypothetical, but will be synthesized in the first amplification cycle) and of priming nucleic acid synthesis from that strand in the direction opposite the first direction and towards the site of hybridization for the first primer. Conditions for conducting such amplifications, particularly under preferred stringent hybridization conditions, are well known. See, for example, PCR Protocols, Cold Spring Harbor Press, 1991.

Other means of producing specific hybridization probes for edg-5 include the cloning of nucleic acid sequences encoding EDG-5 or EDG-5 derivatives into vectors for the production of mRNA probes. Such vectors are known in the art, are commercially available and may be used to synthesize RNA probes in vitro by means of the addition of the appropriate RNA polymerase as T7 or SP6 RNA polymerase and the appropriate reporter

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More particularly, a method for detection of polynucleotides that hybridize with hedg20 7 is exemplified in Example 4, wherein a positive test correlates to approximately at least 70% identity, and more preferably at least 80-85% sequence identity.

It is possible to produce a DNA sequence, or portions thereof, entirely by synthetic chemistry. After synthesis, the nucleic acid sequence can be inserted into any of the many available DNA vectors and their respective host cells using techniques which are well known in the art. Moreover, synthetic chemistry may be used to introduce mutations into the nucleotide sequence. Alternately, a portion of sequence in which a mutation is desired can be synthesized and recombined with longer portion of an existing genomic or recombinant sequence.

The nucleotide sequence for hedg-5 can be used in an assay to detect inflammation or disease associated with abnormal levels of HEDG-5 expression. The cDNA can be labeled by methods known in the art, added to a fluid, cell or tissue sample from a patient, and

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previously defined. fluid is rinsed off, the reporter molecule is quantitated and compared with a standard as with a compatible fluid which optionally contains a reporter molecule. After the compatible incubated under hybridizing conditions. After an incubation period, the sample is washed

NO:12) under stringent conditions. In situ hybridization of chromosomal preparations and invention provides expression products from this locus that hybridize with hedg-5 (SEQ ID using bacterial artificial chromosomes isolated (BACs), as detailed in Example 16. Thus, the for mapping the native gene. The edg-5 gene was mapped to a band p22.3 of chromosome I markers are invaluable in extending genetic maps. Examples of genetic map data can be physical mapping techniques such as linkage analysis using established chromosomal found in the yearly genome issue of Science (e.g. 1994, 265:1981f) The nucleotide sequence for hedg-5 has been used to construct hybridization probes

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investigators searching for disease genes using positional cloning or other gene discovery mapping. The mapping of new genes or nucleotide sequences provide useful landmarks for reveal genes for further investigation. The nucleotide sequence of the subject invention may (Gatti et al (1988) Nature 336:577-580), any sequences mapping to that area may represent or also be used to detect differences in gene sequence between normal and carrier or affected localized by genetic linkage to a particular genomic region, for example, AT to 1 1q22-23 techniques. Once a disease or syndrome, such as ataxia telangiectasia (AT), has been crudely New nucleotide sequences can be assigned to chromosomal subregions by physical

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ä eukaryotic. Host cells may be from the same species from which the nucleotide sequence sequence. The oligopeptide may be expressed in a variety of host cells, either prokaryotic or Gene Expression Technology, Methods and Enzymology, Vol. 185, Academic Press, San polypeptide using well known methods of recombinant DNA technology. Goeddel (1990, was derived or from a different species. Advantages of producing an oligonucleotide by Diego CA) is one among many publications which teach expression of an isolated nucleotide recombinant DNA technology include obtaining adequate amounts of the protein for purification and the availability of simplified purification procedures. Nucleotide sequences encoding edg-5 may be used to produce a purified oligo - or

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5 nucleotides from edg-5 or a desired portion of the polypeptide to a nucleic acid sequence produced from a chimeric nucleotide sequence. This is accomplished by ligating the is more convenient to prepare recombinant proteins in secreted form. Purification steps vary produced by a recombinant cell may be secreted, expressed on cellular membranes, or may be domains and recovery of such peptides from cell culture. EDG-5 (or any of its domains) suitable for the expression of T7Gs, their extracellular, transmembrane or intracellular encoding a polypeptide domain which will facilitate protein purification (Kroll DJ et al with the production process and the particular protein produced. Often an oligopeptide can be contained intracellularly, depending on the particular genetic construction used. In general, it Cells transformed with DNA encoding EDG-5 may be cultured under conditions

ᅜ 20 during direct synthesis and combined with other parts of the peptide using chemical methods direct peptide synthesis using solid-phase techniques (e.g. Stewart at al (1969) Solid-Phase Soc. 85:2149-2154). Automated synthesis may be achieved, for example, using Applied Peptide Synthesis, WH Freeman Co., San Francisco QA; Merrifield J (1963) J Am Chem provided by the manufacturer. Additionally, a particular portion of EDG-5 may be mutated Biosystems 431A Peptide Synthesizer (Foster City, CA) in accordance with the instructions In addition to recombinant production, fragments of EDG-5 may be produced by

(1993) DNA Cell Biol. 12:441-53).

25 mimic a portion of the aa sequence of the protein and may contain the entire aa sequence of a protein must be antigenic. Peptides used to induce specific antibodies may have an aa small naturally occurring molecule such as EDG-5. An antigenic portion of EDG-5 may be sequence consisting of at least five amino acids (aa), preferably at least 10 aa. They should fused to another protein such as keyhole limpel hemocyanin, and the chimeric molecule used for antibody production. EDG-5 for antihody induction does not require biological activity: however, the

is produced against an epitope of the polypeptide and binds to at least part of the natural or recombinant protein. Antibody production includes not only the stimulation of an inunune animal with the polypeptide or an antigenic fragment. An antibody is specific for EDG-5 if it Antibodies specific for EDG-5 may be produced by inoculation of an appropriate

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response by injection into animals, but also analogous processes such as the production of synthetic antibodics, the screening of recombinant immunoglobulin libraries for specific-binding molecules (e.g. Orlandi R et al (1989) PNAS 86:3833-3837, or Huse WD et al (1989) Science 256:1275-1281) or the in vitro stimulation of lymphocyte populations. Current technology (Winter G and MIstein C (1991) Nature 349:293-299) provides for a number of highly specific binding reagents based on the principles of antibody formation. These techniques may be adapted to produce molecules which specifically bind EDG-5s.

An additional embodiment of the subject invention is the use of HEDG-5 specific antibodies, inhibitors, ligands or their analogs as bioactive agents to treat inflammation or disease including, but not limited to viral, bacterial or fungal infections; allergic responses; mechanical injury associated with trauma; hereditary diseases; lymphoma or carcinoma; or other conditions which activate the genes of kidney, lung, heart, lymphoid or tissues of the nervous system.

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Bioactive compositions comprising agonists, antagonists, receptors or inhibitors of HEDG-5 may be administered in a suitable therapeutic dose determined by any of several methodologies including clinical studies on mammalian species to determine maximal tolerable dose and on normal human subjects to determine safe dose. Additionally, the bioactive agent may be complexed with a variety of well established compounds or compositions which enhance stability or pharmacological properties such as half-life. It is contemplated that the therapeutic, bioactive composition may be delivered by intravenous infusion into the bloodstream or any other effective means which could be used for treating problems involving aberrant expression of the Edg-5 gene.

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The examples below are provided to illustrate the subject invention. These examples are provided by way of illustration and are not included for the purpose of limiting the invention.

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EXAMPLES

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Example 1: PCR cloning of murine edg-5 cDNA

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Poly-A+ RNA was isolated from TR and TSM murine neuronal cell lines by twice selecting on oligo-dT cellulose (Pharmacia, Cat. 27-5649-01). 10.5 μg of this RNA was reverse-transcribed with oligo-dT or random hexamers as to prime the RT reaction. RNA and primers were heated to 65°C for 5 min., then cooled to room temperature. Additional reagents were added to give the following final concentrations: 50 mM Tris-Cl, pH 8.3, 6 mM MgCl₂, 40 mM KCl, 1 mM DTT, 1 mM each dNTP, and 1 unit/μ1 of Moloney murine virus RT enzyme.

First strand cDNA was amplified in PCR reactions using degenerate primers A1 (SEQ ID NO: 1) and B1 (SEQ ID NO: 2) was conducted as follows. PCR reactions used 40 ng of first strand cDNA in 10 mM Tris-Cl, pH 8.3, 50 mM KCl, 2 µM of each primer, 1.5 mM MgCl_b, 0.2 µM each dNTP and 2.5 units of Taq DNA polymerase. Thirty pairwise combinations of primers were used. Reactions were placed in a Perkin-Elmer 480 thermal cycler, denatured for 3 min. at 94 °C, and then cycled 25-40 times at 96 °C for 45 sec, 47 °C for 144 sec or 53 °C for 216 sec, and 72 °C for 3 min. initially, increasing 6 sec/cycle. Products were cloned using the TA PCR cloning vector (Invitrogen, Cat. K2000-40). The resulting edg-5 clone, 501(SEQ ID NO: 3) was sequenced by the dideoxy termination method.

20 A1: 5'-AAYTRSATIMTISTIAAYYTIGCIGTIGCIGA-3' (SEQ ID NO: 1)
B1: 5'-CTGIYKWTTCATIAWIMMRTAIAYIAYIGGRTT-3' (SEQ ID NO: 2)

The nucleotide sequence of clone 501(SEQ ID NO: 3) is shown in Figure 1A. A search of Genbank showed that clone 501 (SEQ ID NO: 3) was most closely related to the LPA receptor, also identified in Genbank as the GPCR orphan edg-2 (Genbank MMU70622). Sequence identity between the clone 501 (SEQ ID NO: 3) and edg-2 was 60.5% over the 639 bp length of clone 501 (SEQ ID NO: 3). The amino acid sequence of this nucleotide is shown in Figure 2 (SEQ ID. NO: 15)

Approximately 5x10⁴ phage from an embryonic day 15 whole embryo lambda-ZAP cDNA library (Clontech) was screened with part of this PCR product, 501AB (the original PCR probe from the degenerated PCR screen, using PCR primers A and B), on conventional nylon filter lifts with ³³P-labeled probe, and washed to high stringency. Two clones were

isolated and subcloned into the EcoRI site of pBluescript. One of the clones was sequenced as shown in Figure 1B with the amino acid sequence shown in Figure 5B.

EXAMPLE 2: Isolation of hedg-5 cDNA PCR amplification of partial hedg-5 gene from human genomic DNA

PCR printers JC501-F2 (SEQ ID NO: 4) and JC501-R (SEQ ID NO: 5) were designed using the sequence of clone 501 (SEQ ID NO: 3) and used to obtain a PCR fragment of hedg-5, as detailed below with the ExpandTM PCR system from Boehringer Mannheim (Cat. 1681-842). Human genomic DNA was obtained from Promega (Cat.

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CS01-F2:

15 S'.TTTTTTACTCGAGATTTGCTGGTTATTGCTGTGGAAAG-3' (SEQ ID NO: 4)

CS01-R:

\$'-TTTTTTC'I'AGACGGTCATCACTGTCTTCATTAGCTTC-3' (SEQ ID NO: 5)

20 Each reaction contained the following reagents:

30.25 µl water

10 µl 2.5 mM dNTP mix

5 µl 10x ExpandTM Buffer 3

1.5 µl 10 µM JC501-F2 primer

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1.5 µ1 10 µM JC501-R primer

0.75 μl Expand PCR enzyme (3.5 units/μl)

l μl human genomic DNA (0.272 μg/μl)

PCR Conditions:

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Incubate: 94 ° C for 2 min.

30 cycles: 92°C for 1 min.

45°C for 5 min.

68°C for 1 min.

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Hold:

4°C

On ethidium bromide (EtBr)-stained agarose gel, an intense PCR product of about 390 bp was seen. This product was reamplified in the following PCR reaction:

30.25 µl water

10 μl 2.5 mM dNTP mix

5 μl 10x ExpandTM Buffer 3

1.5 µl 10 µM JC501-F2 primer

1.5 µl 10 µM JC501-R primer

5

0.75 μl Expand PCR enzyme (3.5 units/μl)

1 μl PCR product from the previous PCR reaction

PCR Conditions:

Incubate: 94°C for 2 min.

2

30 cycles: 92°C for 1 min.

45°C for 5 min.

68°C for 1 min.

20

Hold:

4°0

The intense 390 bp product of the PCR reamplification was excised from the agarose gel. The PCR products from 30 µl of the PCR reaction were purified from pooled gel slices using a Qiaquick Gel extraction kit (Qiagen Inc.; Cat. 28706) and eluted with 20 µl of 10 mM Tris-Cl, pH 8.5. The eluted DNA was quantitated and the sequence of the PCR product was determined by automated sequencing at Allelix's in-house facility, with an ABI 377 Sequencer and fluorescent dideoxy terminators, using each primer from the PCR reactions

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Sequencing results showed 81.5% identity at the nucleotide level with the sequence of

mouse clone 501, over a 312 bp overlap excluding the primer sequences.

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PCR amplification and sequencing of large edg-5 cDNA fragments

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to hedg-5, were used to amplify cDNAs encoding larger portions of hedg-5 form a Agt 10 fetal heart cDNA library, as follows. Primers, H501-20F (SEQ ID NO: 6) and H501-246R(SEQ ID NO: 7), specific

H501-20F:

5'-ATGCGGCTGCATAGCAACCTGACCAAAAAG-3' (SEQ ID NO: 6)

H501-246R:

5'-ATCCGCAGGTACACCACAACCATGATGAGG-3' (SEQ ID NO: 7)

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Each reaction contained the following reagents:

30.25 µl water

10 μl 2.5 mM dNTP mix

5 µl 10x Expand M Buffer 3

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1.5 µl 10 µM H501-20F primer

1.5 µl 10 µM H501-246R primer

0.75 μl Expand PCR enzyme (3.5 units/μl)

1 μl fetal heart cDNA library (≥1 library equivalent/μl; Clontech; Cat. HL5017a)

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PCR Conditions:

Incubate: 94°C for 2 min.

30 cycles: 92°C for 1 min.

45°C for 5 min.

68°C for 1.5 min.

25

Incubate: 68°C for 8 min.

Hold: 4°C

30 seen in a fetal heart library, the approximate size expected from the positions of the primers. No specific PCR products were seen in any of 13 other cDNA libraries tested. On EtBr-stained agarose gel, a moderately intense 250 bp PCR product was

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the lgt10 vector, we chose to amplify products only from one direction. The vector-based or JC501-R (SEQ ID NO: 5) primers versus primers derived from the lgt10 vector in which this cDNA library was constructed. Although cDNA inserts are not directionally cloned into the fetal heart cDNA library, PCR reactions were conducted using JC501-F2 (SEQ ID NO: 4) primer sequences were: To obtain additional edg-5 sequence, and possibly amplify the full-length cDNA from

GT10-F: 5'-TTTTGAGCAAGTTCAGCCTGGTTAAGT-3' (SEQ ID NO: 8)

GT10-R: 5'-TGGCTTATGAGTATTTCTTCCAGGGTA-3' (SEQ ID NO: 9)

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GT10-F (SEQ ID NO: 8) vs. JC501-R (SEQ ID NO: 5) primers to amplify the 5' end of edg-5 NO: 9) primers to amplify the 3' end of edg-5 cDNA clones, and another was done with One PCR reaction was done with JC501-F2 (SEQ ID NO: 4) vs. GT10-R (SEQ ID

2 cDNA clones. Each 40 µl reaction contained the following reagents:

23.6 µl water

8.0μ1 2.5 mM dNTP mix

4 μl 10x ExpandTM Buffer 3

2.0 µl 10 µM edg-5 specific primer

20

0.8 μl 10 μM vector primer

0.6 µl Expand PCR enzyme (0.4 units)

Ē cDNA library stock (≥1 library equivalent/µl; Clontech; Cat. HLS017a)

25 PCR Conditions:

Incubate: 94°C for 2 min.

30 cycles: 92 °C for 30 sec

55°C for 2 min.

68°C for 3 min.

Incubate: 68°C for 8 min

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Hold: 4.0

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The results showed 2 faint PCR products (designated 510-5-1 and 510-5-2) from the 3'-end PCR reaction (JC501-F2 (SEQ ID NO:4 /GT10-R (SEQ ID NO:9). From the 5'-end PCR reaction (GT10-I' (SEQ ID NO: 8)/JC501-R (SEQ ID NO:5) again 2 faint PCR bands (designated 510-6-1 and 510-6-2) were seen. Each band was tip-eluted from the gel by stabbing the band with a fresh Pipetman plugged tip, which was then rinsed into 50 µl of TE, pH 8. This solution was used as a stock from which nested reamplifications were done, using the same vector primer vs. a nested human-specific primer as follows:

11.5 µl water

4.0µ1 2.5 mM dNTP mix

5

2 μl 10x ExpandTM Buffer 1

0.6 µl 10 µM edg-5 specific primer

0.6 μl 10 μM vector primer

0.3 µl Expand PCR enzyme (0.4 units)

1 μl tip-eluted PCR DNA stock .

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PCR Conditions:

Incubate: 94°C for 2 min.

30 cycles: 92°C for 30 sec

55 °C for 40 sec

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68°C for 3 min.

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Incubate: 68°C for 8 min.

Hold: 4°C

25 DNA from the most intense band of each nested reamplification was purified using a QlAquick Gel extraction kit and eluted in 50 ul of 10 mM Tris-Cl, pH 8.5.

Full-length cloning of the hedg-5 cDNA into pcDNA3 vector

30 Extension PCR (cycles without primers) was used to extend the overlapping ~1.0 kb 3' fragment (designated 511-5: reamplified from 510-5-2) and 700 bp 5' fragment (designated 511-14: reamplified from 510-6-2) as follows:

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Extension PCR:

19.8µl water

5.6 µl 2.5 mM dNTP mix

4.0 µl 10x Expand Buffer 1

5 μl edg-5 3' PCR DNA fragment (511-5)

5 μl edg-5 5' PCR DNA fragment (511-14)

0.6 μl Expand PCR enzyme (3.5 units/μl)

PCR Conditions:

Incubate 94°C for 2 min.

5

15 cycles: 92°C for 1 min.

60°C for 10 min.

68°C for 3.5 min.

Incubate: 68°C for 8 min

Hold: 4°C

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Two microliters of the extension PCR reaction was then reamplified using the two vector primers (GT10-F (SEQ ID NO:8) and GT10-R (SEQ ID NO:9) to select for full-length

extension products.

32.25 µl water

20

7.0 µl 2.5 mM dNTP mix

5.0 µl 10x ExpandTM Buffer 1

1.5 µl 10 µM GT10-F primer

1.5 µl 10 µM GT10-R primer

0.75 μl Expand PCR enzyme (3.5 units/μl)

25

PCR Conditions:

Incubate 94°C for 2 min.

30 cycles: 92°C for 40 sec 50°C for 40 sec

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68°C for 3 min.

Incubate: 68°C for 8 min.

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Hold: 4.0

Inc., Cat. 28106), cluted in 50 µl of 10 mM Tris-Cl, pH 8.5. The gel-purified PCR fragment was then sent for automated sequencing at Allelix's in-house facility, as described above. The was seen. The PCR product was purified with a QIAquick PCR purification kit (QIAGEN sequencing results confirmed the identity of the amplified band as edg-5, and suggested that a full-length clone of edg-5 had been reconstructed by extension PCR. After gel electrophoresis of the PCR products, an intense DNA band of about 1.4 kb

5 primers GT10-5KXb (SEQ ID NO: 10) and GT10-3BXh (SEQ ID NO: 11). To subclone into pcDNA3 the above DNA was re-amplified with modified vector

GT10-5KXb:

5'-GGGTAGTCGGTACCTCTAGAGCAAGTTCAGCC-3' (SEQ ID NO: 10)

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5'- ATAACAGAGGATCCTCGAGTATTTCTTCCAG-3' (SEQ ID NO: 11)

Reamplification PCR:

67.5 µl water

14 μl 2.5 mM dNTP mix

20

10 µl 10x ExpandTM Buffer 1

3 μl 10 µM GT10-5KXb primer

3 <u>F</u>l 1.5 µl Expand PCR enzyme (3.5 units/µl) 10 µM GT10-3BXh primer

E DNA from previous PCR reaction

25

PCR Conditions:

Incubate 94 °C for 2 min.

5 cycles: 92°C for 1 min.

50°C for 1 min: 68 °C for 2 min.

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25 cycles: 92°C for 1 min.

60°C for 1 min.

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68°C for 2 min.

Incubate: 68°C for 8 min.

Hold:

4. C

pH 8.5 as described previously and restricted with KpnI and XhoI. The PCR product was QIAquick PCR-purified and eluted in 50 µl of 10 mM Tris-Cl.

Two successive restriction digests was performed on the purified extension PCR product as Restriction digest of PCR sample with Kpnl and Xhol:

0 follows:

38 µl Extension PCR DNA

5 <u>F</u>l 10X NEBuffer 1 (New England Biolabs [NEB])

KpnI restriction endonuclease (10 units; NEB, Cat #142S)

10X Acetylated BSA slock (NEB)

2

The restriction digest was incubated for 1 hour in a 37°C water bath, and then the

following reagents and enzyme were added:

10 JL 10X NEBuffer 2 (NEB)

20

Ξ Xhol restriction endonuclease (20 units; NEB, Cat #146S)

Sμl 10X Acetylated BSA stock (NEB)

43 µl water

25 The reaction products were purified using a QIAquick PCR purification kit and cluted in 50

μl of 10 mM Tris-Cl, pH 8.5.

Preparation of pcDNA3 cloning vector with KpnI and Xhol:

4 E pcDNA3 plasmid DNA (Invitrogen; Cat. V790-20) containing a 1.8 kb cDNA

30 insert

10 µl 10X NEBuffer 2 (NEB)

3 µl KpnI restriction endonuclease (NEB: 1:10 dilution; 3 units)

3 µl XhoI restriction endonuclease (NEB: 1:20 dilution; 3 units)

10 µl 10X Acetylated BSA stock (NEB)

64 µl water

S the vector DNA band without cDNA insert was excised, purified using GeneClean II kit (BIO was added and the tubes were incubated for a further 2 hr. The digest was run on a gel, and 101) and eluted in 40 µl of 10 mM Tris-Cl, pH 8.5. The vector DNA was digested for 1 hour at 37°C. Then, 3 units more of each euzyme

5 5 inserts were identified by sequencing, carried out with the in-house ABI automated plasmid vector using T4 DNA ligase kit (NEB, Cat. 202CS) and transformed into Epicurean was plated onto 2xYT/Ampicillin plates and single colonies were picked. DNA minipreps Coli XL-2 Blue MRF' Ultracompetent cells (Stratagene, Cat. 200150). The transformation was chosen for complete sequence determination of the cDNA insert. sequencing system. From this analysis, a clone designated pC3-hedg-5-3 (SEQ ID NO:13) were made using QIAGEN QIA-Prep8 miniprep kit (Cat. 27144) and clones with appropriate The double-digested, gel-purified PCR DNA was ligated into the prepared pcDNA3

Features of the hedg-5 cDNA

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Genbank/EMBL databases (Genbank BTU48236: 55% identity). edg-2, was the highest-scoring full-length cDNA sequence found from the combined sequencing databases indicates that the hedg-5 sequence is novel. The bovine LPA receptor, A BLAST search of Genbank, EMBL, dbEST, and the GSS and STS genomic

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genomic DNA flanking the 5' end of the cDNA sequence (see below). 250 bp of 5' flanking frame of 1059 bp, and a 3'-untranslated region spanning 204 bp. The coding region of edg-5 SEQ ID NO: 12). The proposed translation start site was preceded by an in-frame stop sequence was obtained from a BAC genomic clone as described in Example 16 (Figure 4A 1095-1097. The prediction of this open reading frame is supported by the sequence of begins with the first methionine codon, at nt 36-38 and terminates with the stop codon at nt codon 24 bp upstream. Sequencing of different clones revealed the existence of several This sequence includes 10 bp of 5'-untranslated sequence, the edg-5 open reading

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open reading frame are listed below. Ninc of these substitutions did not result in a change in sequence within the human population. The 15 polymorphisms observed within the edg-5 sequence polymorphisms, which may represent a sampling of natural variability of the edg-5 nonconservative substitutions. the encoded amino acid), while 3 resulted in conservative substitutions and 3 resulted in

Table 1. Apparent polymorphisms in the hedg-5 protein coding region

35					30					23					20					15				10
5 1	917		914		887		874		830		790		788		781		779		716		585		491	Nucleotide Position
GTT	GTC	AAA	AAC	ATT	ATC	GCG	GTG	TTT	TTC	TTT	TCT	TGT	TGC	TIT	TCT	ATT	ATC	GTT	GTC	TTG	CTG	TTT	TTC	Affected Codon & Polymorphism
Valine	Valine	Lysine	Asparagine	Isoleucine	Isoleucine	Alanine	Valine	Phenylalanine	Phenylalanine	Phenylalanine	Serine	Cysteine	Cysteine	Phenylalanine	Serine	Isoleucine	Isoleucine	Valine	Valine	Leucine	Leucine	Phenylalanine	Phenylalanine	Amino Acid Predicted
Silent		Substitution	Conservative	Silent		Substitution	Conservative	Silent		Substitution	Nonconservative	Silent		Substitution	Nonconservative	Silent		Silent		Silent		Silent		Consequence

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922	TCT	Scrine	Nonconservative
	TIT	Phenylalanine	Substitution
1041	CTC	Leucine	Conservative
	ттс	Phenylalanine	Substitution
1277	GAG	Glutamate	

The edg-5 open reading frame of the pC3-hEdg5-3 (SEQ ID NO:13; Figure 3A) clone predicts a 353 amino acid polypeptide (SEQ ID NO: 14, Figure 4A) with many typical features of a GPCR. These include:

GA A

Glutamate

Silent

5

- 1. A hydropathy profile consistent with the 7 transmembrane structure of GPCRs:
- N-terminal extracellular domain: 1-30
- TM-1: 31-56

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- IL-1: 57-63
- TM-2: 64-92
- EL-1: 93-106
- TM-3: 107-125
- IL-2: 126-144

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- TM-4: 145-170
- EL-2: 171-186
- TM-5: 187-207
- IL-3: 208-239
- TM-6: 240-261

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- EL-3: 262-276
- TM-7: 277-297
- C-terminal cytoplasmic domain: 298-353
- 2. Potential N-glycosylation site in the extracellular N-terminal domain, at residue 15
- 30 3. Potential protein kinase C phosphorylation sites at residues 141, 229 and 303
- Potential cAMP- and cGMP-dependent kinase phosphorylation sites at residues 217, 233
- 5. Potential casein kinase-II phosphorylation site at residue 329

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The amino acid sequence of the human edg-5 receptor, SEQ ID NO:14 (Figure 4A), also shows high homology with other members of the edg subfamily of GPCRs. . The pairwise percent identity and similarity is presented in Table 2 below:

5 Table 2.

Percent Amino Acid Identity and Similarity of Edg Family Sequences to the Human Edg-5 receptor

Gene	Percent Identity	Percent Similarity
Ede-1 (Human)	30.1	40.9
Edg-2 (Human)	48.6	59.0
Edg-2 (Bovine)	55.1	
110	77 6	1 LA
Edg-3 (Human)	32.0	40.0
H218 (Edg-4 - Rat)	31.6	40.6
Ede-6 (Human)	46.0	55.5
Tub o (minus)		

Multiple sequence alignment indicates that edg-2 is the closest known relative of edg-5 at the amino acid sequence level, as suggested by the DNA sequence. The edg-5 gene product is also closely related to edg-6, a novel edg gene described in copending application USSN 08/763,938. Edg-2, edg-5 and edg-6 appear to form a subfamily distinct from edg-1, edg-3 and edg-4 within the larger edg gene family.

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8 23 within the N-terminal coding region. The longer open reading frame (Genbank, accession herein, and will produce one or more spliced mRNA variants with a N-terminal extensions. closely with the initiation codon of hedg-5, the edg-5 open reading frame hedg-5 may encode edg-2 and edg-5 and the fact that the methionine codon of the shorter edg-2 product aligns shorter product as amino acid 19 of the longer product. Due to the sequence relatedness of frame (Genbank, accession no:MMU48235), and retains the initiator methionine codon of the no:MMU70622) encodes an 18-amino acid N-terminal extension of the shorter open reading a similar N-terminal extension to the HEDG-5 peptide of SEQ ID NO:14. Such an extension 041) using the approach detailed in start protocols in Molecular Biology (2nd edition, 15-27) 5' RACE using a commercially available 5' RACE kit (Life Technologies, Cat No:18374-Briefly, given the instant disclosure the skilled artisan could discover such splice variants by will result from splicing of sequences found upstream of the hedg-5 sequences presented Briefly, first strand cDNA is primed using an antisense oligonucleotide specific for hedg-5 Alternative splicing variants of murine edg-2 have been found, which differ in length

and ideally directed to a sequence about 500 nucleotides from the 5' end of the known hedg-5 sequence; kidney and lung RNA are preferred templates for cDNA synthesis. Thereafter, first strand cDNA is then tailed using terminal transferase, for example, with deoxyguanine residues. PCR amplification is primed using an anchor primer complementary to the polyguanine tail and a nested primer specific to hedg-5.

EXAMPLE 3: Molecular cloning of hedg5 coding region for expression and functional annlysis in eukaryotic cells.

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unable to obtain a full-length clone. The rarity of edg-5 in cDNA libraries is further supported by a complete absence of EST's from the edg-5 coding regions in the DBEST database, which contains millionsof individual EST's. Therefore, an alternative approach was designed. In this approach, the coding region would be amplified in two fragments from genomic DNA, since we previously determined the location of the single splice site that occurs (between nt 771/772 of SEQUENCE ID NO: 13) in the genomic DNA encoding HEDG5. Then, the two fragments would be joined by an extension PCR in which primers were engineered to contain a 30 bp overlap.between the two fragments to obtain a functional, full-length edg5 cDNA, DNA fragments from two exons next to intron located at nt 996/997 were PCR amplified using the following primers so that they have an overlap of 30 nt.

5' Exon Fragment

HES-261F: [5'-ATGAATGAGTGTCACTATGACAAG-3']
25 HES-1011R: [5'-ATACCACAAACGCCCCTAAGACAGTCATCACCGTCTTC-3']

3' Exon Fragment

HES-982F: [S'-TGATGACTGTCTTAGGGGGCGTTTGTGGTATGCTGGACC-3']

30 HES-1322R: [5'-TTAGGAAGTGCTTTTATTGCAGACTGC-3']

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Human genomic DNA (Clontech, Cat #6550-1) was amplified with each pair of primers under the following condition of PCR amplification by using ExpandTM PCR system from Boehringer-Mannheim (Cat. 1681-842).

Each reaction contained the following reagents:

5.0 μ1 10x PCR Buffer 3
1.0 μ1 25 mM dNTP mix
1.5 μ1 Primer HE5-261F or HE5-982F (10 pmoVI)
1.5 μ1 Primer HE5-1011R or HE5-1322R (10 pmoVI)
0.75 μ1 ExpandTM Enzyme (7.5 units)
38.25 μ1 water
2.0 μ1 DNA

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PCR conditions:

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Incubate: 94°C for 2 min
30 cycles: 94°C for 1 min
55°C for 2 min
68°C for 1 min
68°C for 8 min

Hold:

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DNA fragments of approximately 700 bp (3' exon) and 350 bp (3' exon) were amplified. The two DNA fragments were purified using Qiaquick gel extraction kit (Qiagen, 28.706) and eluted in 50 ul of 10 mM Tris, pH 8.5. Extension PCR (cycles without primers) was then used to join the 5' exon and 3' exon fragments, which overlapped each other by 30 bp.

Extension PCR:

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Each reaction contained the following reagents:

2.0 µl 10x PCR Buffer 3

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0.4 µl 25 mM dNTP mix
                               PCT/CA98/01193
         Hold:
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           გ
                                       PCT/CA98/01193
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0.3 μl Expand^{τM} Enzyme (2.5 units) 13.8 µl water

1.0 µl 3' exon PCR-amplified DNA 1.0 µl 5' exon PCR-amplified DNA

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PCR conditions:

Incubate:

94°C for 2 min

20 cycles: 60°C for 5 min 94°C for 1 min

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68°C for 1.5 min

Hold: Incubate: 68°C for 8 min

∽ Five μl of the amplified product from the above PCR was then reamplified under the following condition of PCR with primers HE5-261F and HE5-1322R, described previously.

Each reaction contained the following reagents:

20 25 35.25 µl 0.75 μ1 1.5 µl Primer HE5-261F (10 pmol/µl) 5.0 µl 10x PCR Buffer 3 1.5 µl Primer HE5-1322R (10 pmol/µl) 1.0 µl 25 mM dNTP mix 5.0 µ1 DNA water ExpandTM Enzyme (7.5 units)

PCR conditions:

25 cycles: Incubate: Incubate: 94°C for 1 min 94°C for 2 min 68°C for 8 min 68°C for 1 min 55°C for 1 min

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sequencing with each primer used in the above PCR reactions, as described previously. The purification kit (Qiagen), cluted in 50 μl of 10 mM Tris, pH 8.5 and was sent for direct PCR resulting sequences showed 93 - 99% identity to human edg5 cDNA, within the edg-5 coding An intense DNA band of about 1.0 kb was purified using the Qiaquick PCR

modified primers HE5-KZKF and HE5-Kpn1322R under the following conditions: To subclone into pcDNA3.1 (Invitrogen; Cat. V795-20) the above DNA was reamplified with

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ت HBS-KZKF: [5'-TTTAAACTCGAGCCACCATGAATGAGTGTCACTATGAC - 3'] HE5-Kpn1322R: [5'-TATATAGGTACCTTAGGAAGTGCTTTTATTGCAGACTGC-

Each reaction contained the following reagents:

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0.75 μΙ $1.0\,\mu l$ 25 mM dNTP mix 5.0 µl 10x PCR Buffer 3 1.5 μl Primer HE5-Kpn1322R (10 pmoVμl) 1.5 µl Primer HE5-KZKF (10 pmoVµl) ExpandTM Enzyme (7.5 units)

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PCR conditions:

Incubate:

94°C for 2 min

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39.25 µl

water

1.0 µ1 DNA

25 cycles: 25 cycles: 94°C for 40 sec 68°C for 1.5 min 50°C for 1 min

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94°C for 40 sec

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68°C for 1.5 min 65°C for 40 sec

Incubate: <u>4</u> 68°C for 8 min

Hold:

Xho1 restriction sites of pcDNA3.1. The PCR product was purified as described previously and subcloned into Kpn1 and

5 EBNA cells together with the 2xSRE-Luciferase reporter plasmid. Plasmid DNA was prepared from several positive clones and cotransfected into 293-

Transient transfection protocol for 293-EBNA:

ょ Day I.

2) NF-kB Reporter Gene Cotransfection: Expression plasmid (3.5 µg) and reporter plasmid 1) 100 mm plates of 293-EBNA with a confluency of ~80% were used for transfection.

DMEM/F12 (serum-free media) and 20 µl Plus Reagent (Lipofectamine Plus Kit, Life (6χNF-kBtk-p4Luc-zeo; 0.5 μg) DNA samples were combined and diluted in 750 μl of

20 Technologies Cat. 10964-013), and incubated at room temperature for 15 min.

3) 30 μ Lipofectamine Reagent (Lipofectamine Plus Kit) was diluted in 750 μ l DMEM/F12. The diluted Lipofectamine was then combined with the DNAPlus mixture and incubated at

4) The 293-EBNA plates were washed once with PBS, and 5 ml DMEM/F12 was added to

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5) DNA/Plus/Lipofectamine mixture was added to each plate of 293-EBNA cells. The plates were left for 3 hr at 37°C in a 5% CO, incubator.

6) The transfection medium was replaced with DMEM/F12 containing 10% FBS to recover

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Day 2.

96-well Blackview plates coated with poly D-lysine (Becton Dickinson Labware, Cat 1) Transfected cells were harvested by trypsinization and 20,000 cells per well were plated in

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wells of the 96-well plate. Cells were returned to the incubator for 24 hr. 40640). Medium was DMEMF12 containing 0.5% FBS. No cells were plated in the outside

1) Media was removed and cells treated with compounds diluted in DMEM/F12 media

containing 0.15% FBS and the following treatments: a) Untreated: DMEM/F12 plus 0.15% FBS; b) AN (10 μM anandamide); c) LPA (10 μM olcyl lysophosphatidic acid).

2) The cells were treated overnight in the incubator.

Day 4.

5 1) Luclite kit (Packard; Cat. 6016911) was used for luciferase assay. All reagents were brought to room temperature before use.

2) Media was removed from each well. 50 µl 0.5M HEPES pH 7.8, 1 mM MgCl₂, 1 mM CaCl, was added to all wells of 96-well plate.

3) Luclite substrate was made up and 50 µl substrate was added to each well as specified in

5 the kit.

4) Plates were incubated at room temperature for 30 min

5) After incubation, plates were counted in a 12-detector Packard Top Count on a program without dark delay.

20 Results:

1998), edg-2, edg-5 and edg-6 proved to be inflammatory LPA receptor subtypes of the edg November 25, 1998 by MUNROE and corresponding PCT application filed on December 30 "Identification of Lysolipid Receptors Involved in Inflammatory Response" filed on As we have documented elsewhere (See U.S. provisional patent application entitled

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receptor family which when activated induce NF-kB. As exemplified in Figure 6, it was determined that HEDG-5, as particularly represented by the two clones pc3-hedg5#3-4 and pc3-hedg5#28, responded to LPA but not anandamide at 10 µM to activate NF-kB. (See

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Three inflammatory subtypes of lysophosphatidic acid (LPA) receptor.

collagenase gene promoter containing an inducible enhancer element for activator protein-l (AP-1) together with the edg-2 and edg-6 receptor sub-types. As shown in Figure 7, the pC3. gene constructs with the serum response element (SRE) or the proximal 1 kb of the human hedg5#28 showed an SRE response and an AP-1 response when treated with 10 µM LPA. An additional experiment was carried out to test the response of clone #28 in reporter

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was cotransfected separately with SRE, NF-kB or AP-1 reporter genes. The AP-1 reporter contained approximately 1 kb of the human collagenase II promoter, and the first 50 bp of the shown to be controlled by AP-1. This transcription factor, like NF-kB has been implicated in transcription: AP-1 and NF-B. Monaldi Arch Chest Dis. 1997 Apr;52(2):178-86. Review.). largely distinct from those of NF-kB (Adcock IM. Transcription factors as activators of gene inflammatory and neoplastic signal transduction., though the gene targets of its action are factor. Cell. 1987 Jun 19;49(6):729-739.), a region whose inducible expression has been inducible genes contain a common cis element recognized by a TPA-modulated trans-acting 5'-untranslated region of the collagenase II transcription unit (Angel P, et al. Phorbol ester-To determine whether these receptors might mediate inflammatory responses, each

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20 cells were pretreated for 6 hr in medium containing 0.5% FBS, then treated overnight in the described above for Example 11, assay #1, except that NF-kB and AP-1 reporter-transfected same medium with or without 10µM LPA. 293-EBNA cells were grown, lipofected in monolayer cultures, and pretreated as

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30 3-4-fold) in the presence of 10 μM LPA, while no response to LPA was seen when the NF. Results: As shown in Fig. 7, all three receptors robustly activated the NF-kB reporter (about edg-5 caused greater than 2-fold induction of the SRE and AP-1 reporters with LPA kB reporter was colransfected with the empty expression vector pcDNA3. With the SRE and transcription through NF-kB, and perhaps, AP-1 as well. Therefore, all three LPA receptors tested here are capable of inducing inflammatory gene untreated control cells). However, edg-6 strongly induced both reporters, while edg-2 and AP-1 reporter genes, some endogenous response to LPA was seen (about 1.5-fold vs

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EXAMPLE 4: Detection of hedg-5 polynucleotides by hybridization with hedg-5.

mutations or through cloning and subsequent manipulations. Moreover, the mammalian nucleotide changes that have accumulated through their divergent evolutionary history. homolog of a given gene usually varies by 10-30% from species to species, as a result of and other highly related genes. Therefore, a method is provided herein for the detection and identification of hedg variants Edg polynucleotides can vary through the introduction of natural or artificial

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ೱ labeling or random priming (several kits are commercially available), or through after agarose gel electrophoresis. The cDNA insert may be labeled using ³³P-nucleotide end-3 or pC3-hedg5#3.4 or pC3-hedg5#28 with appropriate restriction enzymes to release the mixture of polynucleotides are prepared by standard techniques. Examples include Southern known in the art. Nylon filters (e.g. Hybond N+, Cat. RPN132B) bearing a polynucleotide or incorporation of non-natural nucleotides for later detection with antibodies by methods well full-length hedg-5 insert, followed by cDNA insert purification using standard techniques blots, filter lifts from bacterial colonies or bacteriophage plaques and the like. The HEDG-5 coding region of hedg-5 is prepared by restriction of either pC3-hEdg5-

25 Fraction V)], 10% dextran sulfate, 0.1% sodium dodecyl sulfate (SDS)) for 1 hr or more at ml (or enough to cover filters and seal the bag) of hybridization solution (48% deionized solution [50x Denhardt's is 1% Ficoll 400, 1% polyvinylpyrrolidone, 1% BSA (Pentax formamide, 4.8× SSC [20× SSC is 3 M NaCl, 0.3 M sodium citrate, pH 7.0], 1× Denhardt's The dried filters are rehydrated in water, then prehybridized in a scalable bag with 10

30 2 ml of hybridization solution and inject the probe solution into the sealed bag. Sufficient screw-cap tube and incubated in a boiling water bath for 10 min. Transfer the tube to ice, ado volume) at >5×10° cpm/µg DNA. Reseal the bag, mix thoroughly and incubate overnight at probe should be added to give 1 to 15 ng of radiolabeled probe/ml hybridization buffer (final 42°C in a shaking or rotating water bath or incubator. Radiolabeled probe is added to 1 ml of sonicated herring sperm DNA (2 mg/ml) in a

Wash filters three times with 500 ml of low-stringency wash buffer (2× SSC, 0.1% SDS) at RT for 15 min per wash, on a slowly rotating platform. Then wash two times with medium-stringency wash buffer (1× SSC, 0.1% SDS) at 65°C 15 min per wash. Dry the filters and expose to Phosphorimager cassette or autoradiography film. Positive spots or DNA bands are identified after subtraction of background or appropriate negative control samples (see below).

If needed, a DNA spot containing 10 pmol of the full-length hedg insert of pC310 hEdg5-3 can be used as a positive control (Pos) on the filter, and a DNA spot containing 10 pmol of full-length human edg-2 insert (edg-2 open reading frame only) can be used as a negative control (Neg). A full-length open reading frame, or a partial-length open reading frame, of a test DNA (also 10 pmol) will be scored as a positive if the integrated optical density (10D) of the radioactive probe hybridizing to the test DNA (Test) is greater than 15 IOD_{Nex} + (10D_{Pex} – IOD_{Nex})/2. Otherwise, the test DNA will be scored as negative. A positive sequence identity. If a partial-length open reading frame of the test gene is used, then the equivalent regions of edg-5 and edg-2 will be used as positive and negative controls, respectively, for hybridization.

EXAMPLE 5: Antisense analysis

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Knowledge of the correct, complete cDNA sequence of HEDG-5 enables its use as a tool for antisense technology in the investigation of gene function. Oligonucleotides, cDNA or genomic fragments comprising the antisense strand of hedg-5 are used either in vitro or in vivo to inhibit expression of the mRNA. Such technology is now well known in the art, and antisense molecules can be designed at various locations along the nucleotide sequences. By treatment of cells or whole test animals with such antisense sequences, the gene of interest is effectively turned off. Frequently, the function of the gene is ascertained by observing behavior at the intraccllular, cellular, tissue or organismal level (e.g., lethality, loss of differentiated function, changes in morphology, ctc.).

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In addition to using sequences constructed to interrupt transcription of a particular open reading frame, modifications of gene expression is obtained by designing antisense sequences to intron regions, promoter/enhancer elements, or even to trans-acting regulatory genes. Similarly, inhibition is achieved using Hogeboom base-pairing methodology, also

known as "triple helix" base pairing.

EXAMPLE 6: Expression of HEDG-5

Expression of hedg-5 is accomplished by subcloning the cDNAs into appropriate to expression vectors and transfecting the vectors into analogous expression hosts for example E.Coli. In a particular case, the vector is engineered such that it contains a promoter for β-galactosidase, upstream of the cloning site, followed by sequence containing the aminogalactosidase, upstream of the cloning site, followed by sequence containing the aminogalactosidase. Immediately following these terminal Met and the subsequent 7 residues of β-galactosidase. Immediately following these eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage promoter useful for artificial priming and eight residues is an engineered bacteriophage pr

Induction of the isolated, transfected bacterial strain with IPTG using standard methods produces a fusion protein corresponding to the first seven residues of β -methods produces a fusion protein corresponding to the first seven residues of β -methods galactosidase, about 15 residues of "linker", and the peptide encoded within the cDNA. Since galactosidase, about 15 residues of "linker", and the peptide encoded within the cDNA clone inserts are generated by an essentially random process, there is one chance in three that the included cDNA will lie in the correct frame for proper translation. If the cDNA three that the included cDNA will lie in the correct frame for proper translation. If the cDNA is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not in the proper reading frame, and the proper reading frame is not included the proper reading frame in the proper reading frame, it is obtained by deletion or insertion of the appropriate is not included the proper reading frame in the proper reading frame, and the proper reading frame is not included the proper reading frame in the p

25 appropriate length.

The hedg-5 cDNA is shuttled into other vectors known to be useful for expression of protein in specific hosts. Oligonucleotide primers containing cloning sites as well as a segment of DNA (about 25 bases) sufficient to hybridize to stretches at both ends of the target cDNA is synthesized chemically by standard methods. These primers are then used to amplify the desired gene segment by PCR. The resulting gene segment is digested with appropriate restriction enzymes under standard conditions and isolated by gel electrophoresis. Alternately, similar gene segments are produced by digestion of the cDNA with appropriate restriction enzymes. Using appropriate primers, segments of coding sequence from more

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optimize expression by construction of such chimeric sequences. than one gene are ligated together and cloned in appropriate vectors. It is possible to VAO 99/33972

5 manımalian cells such as Chinese Hamster Ovary (CHO) and human 293 cells, insect cells replication to allow propagation in bacteria and a selectable marker such as the β -lactamase such as SM cells, yeast cells such as Saccharomyces cerevisiae, and bacteria such as E. coli include a second selectable marker such as the neomycin phosphotransferase gene to allow antibiotic resistance gene to allow plasmid selection in bacteria. In addition, the vector may For each of these cell systems, a useful expression vector also includes an origin of sclection in transfected eukaryotic host cells. Vectors for use in eukaryotic expression hosts require RNA processing elements such as 3' polyadenylation sequences if such are not part of the cDNA of interest Suitable expression hosts for such chimeric molecules include, but are not limited to,

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5 20 promoters for CHO cells; trp, lac, tac and T7 promoters for bacterial hosts; and alpha factor, expression. Such promoters are host specific and include MMTV, SV40, and metallothionine sarcoma virus enhancer, are used in manimalian host cells. Once homogeneous cultures of alcohol oxidase and PGH promoters for yeast. Transcription enhancers, such as the rous recombinant cells are obtained through standard culture methods, large quantities of Additionally, the vector contains promoters or enhancers which increase

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2 using chromatographic methods known in the art. For example, HEDG-5 can be expressibly recombinantly produced HEDG-5 are recovered from the conditioned medium and analyzed cloned into the expression vector pcDNA3, as exemplified herein. This product can be used for example, using Lipofectamine (Gibco BRL catalog no. 18324-020) mediated gene to transform, for example, HEK293 or COS by methodology standard in the art. Specifically,

EXAMPLE 7: Isolation of Recombinant HEDG-5

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include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules domains added to facilitate protein purification. Such purification facilitating domains that allow purification on immobilized metals, protein A domains that allow purification on immobilized immunoglobulin, and the domain utilized in the FLAGS extension/affinity HEDG-5 is expressed as a chimeric protein with one or more additional polypeptide

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and the HEDG-5 sequence is useful to facilitate expression of HEDG-5. sequence such as Factor XA or enterokinase (Invitrogen) between the purification domain purification system (Immunex Corp., Seattle WA). The inclusion of a cleavable linker

v **EXAMPLE 8: Testing of Chimeric T7Gs**

by Kobilka et al (1988, Science 240:1310-1316) who created a scries of chimeric $\alpha 2\text{-}\beta 2$ transmembrane ligand-receptive sequences of a new isoform with the transmembrane and/or molecule shifted from having more $\alpha 2$ than $\beta 2$ conformation, and intermediate constructs transmembrane sequence into $\beta 2$ -AR. The binding activity of known agonists changed as the adrenergic receptors (AR) by inserting progressively greater amounts of lpha 2-AR intracellular segments of a different T7G for test purposes. This concept was demonstrated demonstrated mixed specificity. The specificity for binding antagonists, however, correlated Functional chimeric T7Gs are constructed by combining the extracellular and/or

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with the source of the domain VII. The importance of T7G domain VII for ligand recognition

domains appears to be preserved throughout the T7G family regardless of category

the yeast receptors are classified as miscellaneous receptors. Thus, functional role of specific was also found in chimeras utilizing two yeast α -factor receptors and is significant because

structural determinants responsible for coupling the receptors to trimeric G-proteins (Dohlman et al (1991) Annu Rev Biochem 60:653-88). A chimeric receptor in which isoform are exchanged with the analogous domains of a known T7G and used to identify the the manner of $\beta 2$ -AR. This demonstrates that for adrenergic-type receptors, G-protein AR was shown to bind ligands with a2-AR specificity, but to stimulate adenylate cyclase in domains V, VI, and the intracellular connecting loop from eta2-AR were substituted into a2corresponding domain on $\beta 2\text{-}AR$ and the resulting receptor bound ligands with $\beta 2\text{-}AR$ was predicted and observed for a chimera in which the V- > VI loop from lphaI-AR replaced the recognition is present in domains V and VI and their connecting loop. The opposite situation In parallel fashion, internal segments or cytoplasmic domains from a particular

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엉 specificity and activated G-protein-mediated phosphatidylinositol turnover in the α 1-AR manner. Finally, chimeras constructed from muscarinic receptors also demonstrated that V-> VI loop is the major determinant for specificity of G-protein activity (Bolander FF, supra).

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Chimeric or modified T7Gs containing substitutions in the extracellular and transmembrane regions have shown that these portions of the receptor determine ligand binding specificity. For example, two Ser residues conserved in domain V of all adrenergic and D catecholamine T7G teceptors are necessary for potent agonist activity. These serines are helieved to form hydrogen bonds with the catechol moiety of the agonists within the T7G binding site. Similarly, an Asp residue present in domain III of all T7Gs which bind biogenic amines is believed to form an ion pair with the ligand amine group in the T7G binding site.

Functional, cloned T7Gs are expressed in heterologous expression systems and their biological activity assessed (e.g. Marullo et al (1988) Proc Natl Acad Sci 85:7551-55; King et al (1990) Science 250:121-23). One heterologous system introduces genes for a mammalian T7G and a mammalian G-protein into yeast cells. The T7G is shown to have appropriate ligand specificity and affinity and trigger appropriate biological activation—growth arrest and morphological changes—of the yeast cells.

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An alternate procedure for testing chimeric receptors is based on the procedure utilizing the P₂₀ purinergic receptor (P₂₀) as published by Erb et al (1993, Proc Natl Acad Sci 90:104411-53). Function is easily tested in cultured K562 human leukemia cells because these cells lack P₂₀ receptors. K562 cells are transfected with expression vectors containing either normal or chimeric P₂₀ and loaded with fura-a, fluorescent probe for Ca++. Activation of properly assembled and functional P₂₀ receptors with extracellular UTP or ATP mobilizes intracellular Ca++ which reacts with fura-a and is measured spectrofluorometrically. As with the T7G receptors above, chimeric genes are created by combining sequences for extracellular receptive segments of any newly discovered T7G polypeptide with the nucleotides for the transmembrane and intracellular segments of the known P₂₀ molecule. Bathing the transfected K562 cells in microwells containing appropriate ligands triggers binding and fluorescent activity defining effectors of the T7G molecule. Once ligand and function are established, the P₂₀ system is useful for defining antagonists or inhibitors which block binding and prevent such fluorescent reactions.

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EXAMPLE 9: Production of HEDG-5 Specific Antibodies

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Two approaches are utilized to raise antibodies to HEDG-5, and each approach is useful for generating either polyclonal or monoclonal antibodies. In one approach, denatured

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protein from reverse phase HPLC separation is obtained in quantities up to 75 mg. This denatured protein is used to immunize mice or rabbits using standard protecols; about 100 micrograms are adequate for immunization of a mouse, while up to 1 mg might be used to immunize a rabbit. For identifying mouse hybridomas, the denatured protein is radioiodinated and used to screen potential murine B-cell hybridomas for those which that 20 m

produce antibody. This procedure requires only small quantities of protein, such that 20 mg is sufficient for labeling and screening of several thousand clones.

In the second approach, the amino acid sequence of an appropriate HEDG-5 domain, as deduced from translation of the cDNA, is analyzed to determine regions of high

antigenicity. Oligopeptides comprising appropriate hydrophilic regions are synthesized and antigenicity. Oligopeptides comprising appropriate hydrophilic regions are synthesized and used in suitable inumunization protocols to raise antibodies. Analysis to select appropriate used in suitable inumunization protocols to raise antibodies. Analysis to select appropriate epitopes is described by Ausubel FM et al (supra). The optimal amino acid sequences for immunization are usually at the C-terminus, the N-terminus and those intervening, immunization are usually at the C-terminus, the N-terminus and those intervening.

15 hydrophilic regions of the polypeptide which are likely to be exposed to the external environment when the protein is in its natural conformation.

Typically, selected peptides, about 15 residues in length, are synthesized using an Applied Biosystems Peptide Synthesizer Model 431A using fmoc-chemistry and coupled to keyhole limpet hemocyanin (KLH; Sigma, St. Louis MO) by reaction with M-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS; Ausubel FM et al, supra). If necessary, a cysteine is introduced at the N-terminus of the peptide to permit coupling to KLH. Rabbits are immunized with the peptide-KLH complex in complete Freund's adjuvant. The resulting antisera are tested for antipeptide activity by binding the peptide to plastic, blocking with 1% antisera are tested for antipeptide activity by binding and reacting with labeled (radioactive bovine sewm albumin, reacting with antisera, washing and reacting with labeled (radioactive or fluorescent), affinity purified, specific goal anti-rabbit lgG.

Hybridomas are prepared and screened using standard techniques. Hybridomas of interest are detected by screening with labeled HEDG-5 to identify those fusions producing the monoclonal antibody with the desired specificity. In a typical protocol, wells of plates (FAST; Becton-Dickinson, Palo Alto CA) are coated during incubation with affinity purified, specific rabbit anti-mouse (or suitable antispecies lg) antibodies at 10 mg/ml. The coated wells are blocked with 1% BSA, washed and incubated with supernatants from hybridomas.

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After washing the wells are incubated with labeled HEDG-5 at 1 mg/ml. Supernatants with specific antibodics bind more labeled HEDG-5 than is detectable in the background. Then clones producing specific antibodies are expanded and subjected to two cycles of cloning at limiting dilution. Cloned hybridomas are injected into pristane-treated mice to produce ascites, and monoclonal antibody is purified from mouse ascetic fluid by affinity chromatography on Protein A. Monoclonal antibodies with affinities of at least 10⁴ M⁻¹, preferably 10° to 10⁴⁰ or stronger, are typically made by standard procedures as described in Harlow and Lane (1988) Antihodies: A Laboratory Manual, Cold Spring Harbor Laboratory.

EXAMPLE 10: Diagnostic Test Using HEDG-5 Specific Antibodies

Practice, Academic Press, New York City, both incorporated herein by reference.

Cold Spring Harbor, NY; and in Goding (1986) Monoclonal Antibodies: Principles and

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Particular HEDG-5 antibodies are useful for investigating signal transduction and the diagnosis of infectious or hereditary conditions which are characterized by differences in the amount or distribution of HEDG-5 or downstream products of an active signaling cascade.

Diagnostic tests for HEDG-5 include methods utilizing antibody and a label to detect HEDG-5 in human body fluids, membranes, cells, tissues or extracts of such. The

polypeptides and antibodies of the present invention are used with or without modification.

Frequently, the polypeptides and antibodies are labeled by joining them, either covalently or noncovalently, with a substance which provides for a detectable signal. A wide variety of labels and conjugation techniques are known and have been reported extensively in both the scientific and patent literature. Suitable labels include radionuclides, enzymes, substrates, cofactors, inhibitors, fluorescent agents, chemiluminescent agents, chromogenic agents, magnetic particles and the like. Patents teaching the use of such labels include US Patent No's. 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149; and 4,366,241.

Also, recombinant immunoglobulins may be produced as shown in US Patent No.4,816,567, Incorporated herein by reference.

A variety of protocols for measuring soluble or membrane-bound HEDG-5, using either polyclonal or monoclonal antibodies specific for the protein, are known in the art. Examples include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA) and fluorescent activated cell sorting (FACS). A two-site monoclonal-based immunoassay

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utilizing monoclonal antibodies reactive to two non-interfering epitopes on HEDG-5 is preferred, but a competitive binding assay may be employed. These assays are described, among other places, in Maddox, DE et al (1983, J Exp. Med. 158:1211f).

EXAMPLE 11: Purification of Native HEDG-5 Using Specific Antibodies

Native or recombinant HEDG-5 is purified by immunoaffinity chromatography using antibodies specific for HEDG-5. In general, an immunoaffinity column is constructed by covalently coupling the anti-TRH antibody to an activated chromatographic resin.

Polyclonal immunoglobulins are prepared from immune sera either by precipitation with ammonium sulfate or by purification on immobilized Protein A (Pharmacia LKB Biotechnology, Piscataway NJ). Likewise, monoclonal antibodies are prepared from mouse ascites fluid by ammonium sulfate precipitation or chromatography on immobilized Protein A. Partially purified immunoglobulin is covalently attached to a chromatographic resin such as CnBr-activated Sepharose (Pharmacia LKB Biotechnology). The antibody is coupled to the resin, the resin is blocked, and the derivative resin is washed according to the manufacturer's instructions.

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20 Such immunoaffinity columns are utilized in the purification of HEDG-5 by preparing a fraction from cells containing HEDG-5 in a soluble form. This preparation is derived by solubilization of whole cells or of a subcellular fraction obtained via differential centrifugation (with or without addition of detergent) or by other methods well known in the art. Alternatively, soluble HEDG-5 containing a signal sequence is secreted in useful quantity into the medium in which the cells are grown.

A soluble HEDG-5-containing preparation is passed over the immunoaffinity column, and the column is washed under conditions that allow the preferential absorbance of HEDG-5 (e.g., high ionic strength buffers in the presence of detergent). Then, the column is eluted under conditions that disrupt antibody/protein binding (e.g., a buffer of pH 2-3 or a high concentration of a chaotrope such as urea or thiocyanate ion), and HEDG-5 is collected.

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EXAMPLE 12: Drug Screening

coupled receptor such as EDG-5 can be measured using any of a variety of appropriate potentially used to identify HEDG-5 ligands. For example, the activity of a G protein HEDG-5 is a G protein coupled receptor any of the methods commonly used in the art may mobilization, or inositol phospholipid hydrolysis. More particularly, activation of EDG-5 can detectable gene product, for example, CAT or luciferase. Alternatively, the cell is loaded approach, measures the effect of ligand binding on the activation of intracellular second be measured using the NF-kB, SRE and/or AP-1 functional assays, as described above. One level of some second messenger system, such as adenylate cyclase, guanylyl cyclase, calcium functional assays in which activation of the receptor results in an observable change in the HEDG-5 or binding fragments thereof in any of a variety of drug screening techniques. As calcium indicate modulation of the receptor as a result of ligand binding. Thus, the present with a reporter substance, e.g., FURA whereby changes in the intracellular concentration of which is sensitive to the level of that second messenger controlling expression of an easily messenger pathways, using a reporter gene. Typically, the reporter gene will have a promoter invention provides methods of screening for drugs or any other agents which affect signal This invention is particularly useful for screening therapeutic compounds by using

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Alternatively, the polypeptide or fragment employed in such a test is either free in solution, affixed to a solid support, borne on a cell surface or located intracellularly. One method of drug screening utilizes eukaryotic or prokaryotic host cells (or membrane preparations therefrom) which are stably transformed with recombinant nucleic acids competition binding assays. P-labelled LPA could be used in such a competition binding assays. YP-labelled LPA could be used in such a competition binding assays. One measures, for example, the formation of complexes between HEDG-5 and the agent being tested. Alternatively, one examines the diminution in complex formation between HEDG-5 and a ligand, for example LPA, caused by the agent being tested.

EXAMPLE 13: Rational Drug Design

Herein, the goal of rational drug design is to produce structural analogs of biologically active phospholipids of interest or of small molecules with which they interact, agonists,

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antagonists, or inhibitors. Any of these examples are used to fashion drugs which are more active or stable forms of the phospholipid or which enhance or interfere with the function of a phospholipid in vivo.

protein-inhibitor complex, is determined by x-ray crystallography, by computer modeling or, most typically, by a combination of the two approaches. Both the shape and charges of the polypeptide must be ascertained to elucidate the structure and to determine active site(s) of the molecule. Less often, useful information regarding the structure of a polypeptide is gained by modeling based on the structure of homologous proteins. In both cases, relevant structural information is used to design efficient inhibitors. Useful examples of rational drug design includes molecules which have improved activity or stability as shown by Braxton S and Wells JA (1992, Biochemistry 31:7796-7801) or which act as inhibitors, agonists, or antagonists of native peptides as shown by Athauda SB et al (1993 J Biochem 113:742-46), incorporated herein by reference.

EXAMPLE 14: Use and Administration of Antibodics, Inhibitors, or Antagonists

Antibodies, inhibitors, or antagonists of HEDG-5 (or other treatments to limit signal transduction, LST) provide different effects when administered therapeutically. LSTs are formulated in a nontoxic, inerl, pharmaceutically acceptable aqueous carrier medium preferably at a pH of about 5 to 8, more preferably 6 to 8, although pH may vary according to the characteristics of the antibody, inhibitor, or antagonist being formulated and the condition to be treated. Characteristics of LSTs include solubility of the molecule, half-life and antigenicity/immunogenicity. These and other characteristics aid in defining an effective

topical creams and gels; transmucosal spray and aerosol; transdermal patch and bandage;

30 injectable, intravenous and lavage formulations; and orally administered liquids and pills

particularly formulated to resist stomach acid and enzymes. The particular formulation, exact

dosage, and route of administration is determined by the attending physician and varies

according to each specific situation.

LSTs are delivered by known routes of administration including but not limited to

Such determinations are made by considering multiple variables such as the condition to be treated, the LST to be administered, and the pharmacokinetic profile of a particular LST. Additional factors which are taken into account include severity of the disease state, patient's age, weight, gender and diet, time and frequency of LST administration, possible combination with other drugs, reaction sensitivities, and tolerance/response to therapy. Long acting LST formulations might be administered every 3 to 4 days, every week, or once every two weeks depending on half-life and clearance rate of the particular LST.

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Normal dosage amounts vary from 0.1 to 100,000 micrograms, up to a total dose of about 1 g, depending upon the route of administration. Guidance as to particular dosages and methods of delivery is provided in the literature; see US Patent Nos. 4,657,760; 5,206,344; or 5,225,212. Those skilled in the art employ different formulations for different LSTs. Administration to cells such as nerve cells necessitates delivery in a manner different from that to other cells such as vascular endothelial cells.

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It is contemplated that abnormal signal transduction, trauma, or diseases which trigger HEDG-5 activity are treatable with LSTs. These conditions or diseases are specifically diagnosed by the tests discussed above, and such testing should be performed in suspected cases of viral, bacterial or fungal infections: allergic responses; mechanical injury associated with trauma; hereditary diseases; lymphoma or carcinoma; or other conditions which activate the genes of lymphoid or neuronal tissues.

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EXAMPLE 15: Production of Transgenic Animals

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Animal model systems which elucidate the physiological and behavioral roles of the HEDG-5 receptor are produced by creating transgenic animals in which the activity of the HEDG-5 receptor is either increased or decreased, or the amino acid sequence of the expressed HEDG-5 receptor is altered, by a variety of techniques. Examples of these techniques include, but are not limited to: 1) Insertion of normal or mutant versions of DNA encoding a HEDG-5 receptor, by microinjection, electroporation, retroviral transfection or other means well known to those skilled in the art, into appropriate fertilized embryos in order to produce a transgenic animal or 2) Homologous recombination of mutant or normal, human or animal versions of these genes with the native gene locus in transgenic animals to alter the regulation of expression or the structure of these HEDG-5 receptor sequences. The

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technique of homologous recombination is well known in the art. It replaces the native gene with the inserted gene and so is useful for producing an animal that cannot express native HEDG-5 receptors but does express, for example, an inserted mutant HEDG-5 receptor, which has replaced the native HEDG-5 receptor in the animal's genome by recombination, resulting in underexpression of the transporter. Microinjection adds genes to the genome, but does not remove them, and so is useful for producing an animal which expresses its own and added HEDG-5 receptors, resulting in overexpression of the HEDG-5 receptors.

5 20 5 is as follows: Female mice are mated, and the resulting fertilized eggs are dissected out of tissue specific regulatory elements may be fused with the coding region to permit tissuetheir oviducts. The eggs are stored in an appropriate medium such as M2 medium. DNA or specific expression of the trans-gene. The DNA, in an appropriately buffered solution, is put experimental means to regulate expression of the transgene. Alternatively or in addition, Inducible promoters may be fused with the coding region of the DNA to provide an cDNA encoding a HEDG-5 purified from a vector by methods well known in the art. into the oviduct of a pseudopregnant mouse (a mouse stimulated by the appropriate pronucleus of the egg, and the DNA solution is injected. The injected egg is then transferred and the egg to be injected is put in a depression slide. The needle is inserted into the into a microinjection needle (which may be made from capillary tubing using a piper puller) uterus, implants, and develops to term. As noted above, microinjection is not the only hormones to maintain pregnancy but which is not actually pregnant), where it proceeds to the methods for inserting DNA into the egg cell, and is used here only for exemplary purposes. One means available for producing a transgenic animal, with a mouse as an example

25 EXAMPLE 16: Isolation, Chromosomal Localization and Partial Sequencing of a hedg-5 Genomic Clone

To identify genomic clones containing the hedg-5 gene, the H501-20F (SEQ ID NO: 6) and H501-246R (SEQ ID NO: 7) primers were used to amplify human genomic DNA as 30 described in Example 2. One microliter of human genomic DNA (Clontech; Cat #6550-1) was used as template. The PCR product was purified and sequenced in-house, using the PCR primers to prime the sequencing reactions. The sequence of this product (see SEQ ID. NO: 12) matched the cDNA sequence previously obtained for hedg-5 (see SEQ ID. NO: 13),

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indicating that these primers could be used to identify genomic clones containing this region of the hedg-5 gene

clones represent at least part of the hedg-5 gene. The BAC-28 (IF) clone was subsequently clones were identified by this method: BAC-28 (IF) and BAC-236 (I3M). Once the DNA haploid genome equivalents were screened using the edg-5 diagnostic PCR primers. Two constructs with ~120 kb human genomic DNA inserts. In total, clones representing about 3 PCR using these primers. The library contained bacterial artificial chromosome (BAC) at Genome Systems Inc. The locus for the hedg-5 gene mapped to band p22.3 of human from these clones was received, their identity was verified in-house by sequencing of the used to localize the gene on human chromosomes by fluorescent in situ hybridization (FISH) PCR product we obtained using the edg-5 diagnostic primers: this analysis showed both An arrayed library of genomic DNA clones (Genome Sciences Inc.) was screened by

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20 oncogenes), while the second is an inherited syndrome in which the sixth and seventh cranial suppresser of cellular transformation (a class of genes called tumor suppressers or anti-Suppression-1 or MTS1) and 157900 (Moebius Syndrome). The first represents a dominant been cloned. These were the database entries 154280 (Malignant Transformation for inherited diseases which genetically map to this region, but for which genes have not yet to either of these phenotypes is not known nerves are small or absent, leading to facial paralysis. Whether edg-5 gene defects contribute A search of the on-line Mendelian Inheritance in Man database revealed two entries

hedg-55, revealing 250 bp of genomic DNA sequence upstream of the 5' end of the cDNA by the arrowhead between nt 996/997 of the sequence shown in Figure 4A. This intron falls showed that only one intron exists within the coding region of hedg-5, at a position indicated position(s) of introns (if any) within the coding region of the edg-5 gene. Sequencing results performed in the region flanking the 5' end of the edg-5 cDNA sequence derived from pC3. within the codon for Gly-246 of the edg-5 amino acid sequence. Additional sequencing was Sequencing was performed on DNA prepared from BAC-28 (1F) to determine the

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EXAMPLE 17: Expression and tissue distribution of Edg-5 RNA in the rat.

5 spleen and liver. Muscle tissue may also express low levels of edg-5 mRNA. Further, anti-1007) were prohed with radiolabeled edg-5 cDNA. Washing was performed at high expression levels are highest in lung, kidney and testis. Lower RNA levels were seen in skin blots were then subjected to autoradiography. The Northern blot results show that RNA stringency conditions that do not permit detection of edg-2 or other related transcripts. The known in the art. Two different multi-tissue rat RNA blots (Origene .Cat. MB-1005 and MBheart, small intestine and stomach. Little or no delectable RNA was found in thymus, brain Northern blotting was carried out with the edg-5 cDNA insert by techniques well-

2 invention will be apparent to those skilled in the art without departing from the scope and preferred embodiments, it should be understood that the invention as claimed should not be spirit of the invention. Although the invention has been described in connection with specific unduly limited to such specific embodiments. Various modifications and variations of the described method and system of the

those of skill in the art to for in situ hybridization expression studies.

sense oligonucletide probes based on the hedg-5 sequence disclosed herein can be used by

CLAIMS

- An isolated nucleotide sequence encoding a mammalian EDG-5 receptor or biologically active fragment thereof.
- The isolated nucleotide sequence of claim 1 encoding a murine EDG-5 receptor or biologically active fragment thereof.
- The isolated nucleotide sequence of claim 2 encoding a murine EDG-5 receptor of
- 10 Figure 1B or biologically active fragment thereof.
- 4. The isolated nucleotide sequence of claim 1 encoding a human EDG-5 receptor or biologically active fragment thereof.
- 15 5. The isolated nucleotide sequence of claim 1 wherein the biologically active fragment is activated by LPA.
- An isolated nucleotide sequence selected from the group consisting of:
- (a) the nucleotide sequence comprising nucleotides 36-1907 of SEQ. ID NO: 12
- (b) the nucleotide sequence of Figure 3B;

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- (c) the nucleotide sequence of Figure 3C;
- (d) a nucleotide sequence comprising at least about 70% sequence identity to (a), (b)
- or (c) and which hybridizes under stringent conditions to the nucleotide sequence of (a), (b) or (c), respectively; and
- 25 (e) the nucleotide sequence which encodes the amino acid sequence of Figure 4A, 4B or 4C.
- The isolated nucleotide sequence of Claim 6 wherein the nucleotide sequence is selected from the group consisting of:
- 30 (1) the nucleotide sequence of (a), (b), (c) or (e) of claim 6; and
- (2) the nucleotide sequence of (d) of claim 6 wherein the nucleotide sequence has at least about 80-85% sequence identity to the nucleotide sequence of (a), (b) or (c) of claim 6.

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- 8. The isolated nucleotide sequence of Claim 6 wherein the nucleotide sequence is selected the group consisting of:
- (1) the nucleotide sequence of (a), (b), (c) or (e) of claim 6; and
- (2) the nucleotide sequence of (d) of claim 6 wherein the nucleotide sequence has at
- least about 95% sequence identity to the nucleotide sequence of (a), (b) or (c) of claim

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The complement of the nucleotide sequence of Claim 8.

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- 10 10. An expression vector comprising the nucleotide sequence of Claim 8.
- 11. A host cell comprising the expression vector of Claim 10.
- 12. The isolated and purified amino acid sequence for the HEDG-5 receptor encoded by
- 15 the nucleotide sequence of claim 8.
- 13. The isolated and purified amino acid sequence of claim 12 comprising the amino acid sequence of SEQ. ID NO:13 (Figure 4A), Figure 4B or Figure 4C or a biological active portion thereof.
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- 14. The isolated nucleotide sequence of Claim 6 wherein the nucleotide sequence is selected from the group consisting of the nucleotide sequence which encodes the amino acid sequence of SEQ ID NO: 13 (Figure 4A), Figure 4B and Figure 4C.
- 25 15. A hybridization probe of the nucleotide sequence of Claim 5.
- 16. A method of screening compounds to identify HEDG-5 ligands comprising the steps
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- (a) culturing cells which express the HEDG-5 receptor or with a membrane
- preparation obtained therefrom; and

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(b) contacting said compound with said cells or said membrane preparation; and (c) determining whether binding between the HEDG-5 receptor and the candidate ligand has occurred.

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A HEDG-5 ligand identified by the method of claim 16.

A method of screening compounds to identify HEDG-5 antagonists comprising the

- culturing cells which express the HEDG-5 receptor or with a membrane
- an agonist and said compound to be tested for antagonist activity at said receptor; and preparation obtained therefrom; contacting said cells or said membrane preparation with a mixture comprising
- agonist and said receptor absent the antagonist. comparing this measured response with a standard response for binding between said indicative of the degree of binding between said agonist and said receptor and determining the degree of antagonist activity by measuring a response

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- The method of claim 18 wherein said agonist is LPA.
- 20. An antagonist identified by the method of claim 18.

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ABSTRACT

homologs. The invention also provides methods for determining agonists and antagonists for for mammalian EDG-5 receptor homologs, and particularly for human EDG-5 receptor The present invention is directed to nucleic acid sequences and amino acid sequences

agonists for EDG-5 receptors. disorders associated with aberrant expression or activity of EDG-5. SIP and SPC are EDG-5 receptors in addition to assays, expression vectors, host cells and methods for treating

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Figure 1A: DNA sequence of the murine edg-5 RT-PCR clone 501 (SEQ ID NO: 3)

100 251 201 151 AACTTGACCA AAAAGCGGGT GACGCTGCTC ATTCTGCTGG TGTGGGCCAT 101 TTATTGCTGT GGAAAGACAC ATGTCNATCA TGAGGATGAG AGTCCACAGC 451 101 351 CCAGGGGCTC CTAGACACCA GCCTGACTGC CTCCCTGGCC AATTTGCTGG AACACTGGCC CGGTGTCGAA AACGTTGACC GTCAACCGCT GGTTCCTCCG AACACGTGAA GNGCTGGTTC CTGCTGCTCG CACTGCTCAA CTCCGTCATG ATACGTACGC ATCTACATGT ATGTTANAAG GAAAACCAAC GTCTTATCTC ACATCTCGGC CTGCTCTTCT CTGGCTCCCA TTTACAGTAG GAGTTACCTC COCCATCITC ATGGGGGCCG TCCCCACNCT GGGATGGAAT TGCCTCTGCA AACCCCCTCA TCTACTGCCG CTCTCCNNAC TTTCCATGG AAGACAGTGA TGACCOTCTT AGGCGCCTTC GTGGTGTGCT GGACCCCGGG CÁCACACCAG TGGCTCCATC AGCCGCCGGA GGGCTCCCAT GAAGCTAATG ATTITCIDGA CIGITICCAA CCICCIGGCC TICITCATCA IGGIGGCGGI TETGGTGGTT CTGCTGCTGG ACGGCCTGAA CTGCAAGCAG TGTAACGTGC

Figure 1B: Sequence of full-length mEDG-5 cDNA insert and alignment with MEDG-5 amino acid sequence. Translation starts at nt 19. Translation termination codon is located at nt

241 181 121 303 13 M N E C H Y D K R M D F F Y
GCACAGTTCTTGTCCACCATGAATGAGTGTCACTATGACAAGGGCATGGACTTTTTCTAC C V G T P F C L P I F F S N S L V I A A TGCGTGGGGACGTTCTTCTGCCTCTTTATATTTTTTTCTAACTCCCTGGTCATTGCTGCG K T L T V N R W F L R Q G L L D T S L T
AAAACGTTGACCGCTCAACCGCTGGTTCCTCCGCCAGGGGCTCCTAGACACCAGCCTGACT A D P P A G I A Y V P L M P N T G P V S GCGGATTTCTTCGCCGGAATCGCTTACGTGTTCCTGATGTTTAACACTGGCCCGGTGTCG VITHRKPHFPFYYLLANLAA AACAGGAGCAACACAGACACAGCGGACGACGAGTGGACAAGGGTAGAAGCTTGTGATCGTCCTG **GTGATCACAAACCGGAAGTTCCACTTTCCCTTCTACTACCTGCTGGCTAACTTAGCTGCT** RSNIDTADBWIGIKLVIVL 180 120 60 360 300 240

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Figure 1B (cont.)

GTCTCACAAAGCACGTGGACAAGGGTTGTTTGAGGGCTCCATGCATCACTTCTGGGGCTTT	TANGCCACGGACGCCTCCGCCCTCTTCCCCCTGGGGAAAGAGCTGTTAAGCGTCCTCACCT	G S Q Y L S D S I S Q G P V C N K N G S GGCAGCCAGTACCTGGAGGACAAGCATCAGCCCAGGCCCGGTGTGCAATAAAAACGGCTCC 1021	S N T E R R P S R N P S T I H S R S E T AGCAATACCGAGAGGGCGCCCCCCCCCCACCATCCACCAGCAGGAGCGAGACG 961	Y K D E D M Y N T H R K H I C C A L Q D TACAAGGACGAGGACATGTACAACACCATGCGGAAGATGATCTGCTGTGCCCTGCAGGAC 901	KRWPLLLALLNSVMNPIIYS AAGCGTTGGTTCTGCTCTGCACTCATCTATCTACTCTATCTA	G L V V L L L D G L N C K Q C N V Q H V GOTCTGGTGGTTCTGCTGCTGGACGGCCTGAACTGCAAGCAGTGTAACGTGCAACACGTG	M K L M K T V M T V L G A F V V C W T P ATGAAGCTAATGAAGAGATGATGACGGTCTTAGGCGCCTTCGTGGTGTGCTGGACCCCC 721	R K T N V L S P H T S G S I S R R R A P AGGAMANCCAACGTCTTATCTCCACCACACACGAGGGCTCCXTCAGCCGCGGAGGGCTCCC	N L L A P F I M V A V Y V R I Y M Y V K AACCTCCTGGCCTTCTTCATCATGTGGCGGTATACGTACG	A C S S L A P I Y S R S Y L I F H T V S GCCTGCTCTTCTCTGGGCTCCCATTTACAGTAGGAGTTACCTCATTTTCTGGACTGTGTCC 541	I A I P M G A V P T L G M N C L C N I S ATCGCCATCTTCATGGGGCCGTCCCCACGCTGGGATGGAATTGCCTCTGCAACATCTCG	R V H S N L T K K R V T L L I L L V H A AGAGTCCACAGCAACTTGACCGAAAAAGCGGGTGACGCTGATCATTCTGCTGGTGTGGGCC	A S L A N L L V I A V B R H M S I M R H GCCTCCCTGGCCAATTGCTGGTTATTGCTGTGGAAAGACACATGTCAATCATGAGGATG 361
GGCTCCATGCATCACTTCTGGGGCTTT	CACGGACGCCTCCGCCCTCTTCCCCCTGGGGAAAGAGCTGTTAAGCGTCCTCACCT	G P V C N K N G S GGCCCGGTGTGCAATAAAACGGCTCC	S T I H S R S E T TTCCACCATCCACAGCAGAGCGAGACG	E D M Y N T M R K M I C C A L Q D COARGARCATOTRACAACACTOCOGAAAGATGATCTGCTGTGCCCTGCAGGAC	LALLNSVMNPITYS GCTCGCACTGCTCACTCCGTCATCTACTCG	C K Q C N V Q H V TGCAAGCAGTGTAACGTGCAACACGTG	Q A F V V C H T P NGCCCTTCGTGGTGTGCTGGACCCCG	G S I S R R R A P	Y V R I Y M Y V K ATROGRACOCATOTACATOTATAAA	S Y L I F H T V S AGTTACCTCATTTTCTGGACTGTGTCC	G W N C L C N I S GGATGGAATTGCCTCTGCAACATCTCG	T L L I L L V W A ACCCTACTCATTCTCCTGGTGTGGGCC	B R H M S I M R H GAAAGACACATGTCAATCATGAGGATG
1200	1140	1080	1020	960	900	840	780	720	660	600	540	8 0	420

Pigure 1B (cont.)

1201 1361 1561 1441 1681 1621 1861 1001 1741 1501 1921 1981 TAAGTTTTCATGGTCAAGGAAAATAGATTTACGGCGTTTAGTAAAGCGCACAGGAAAAGG CGAGAGCATGCTGGGCTCTGTCACCATCTTGCCACCATTGTCTTGTGTGTTTTTCAATGATG AGAGATGAGCAGTGGGTTCCGGCTTGTCTGTGATCCGCTCCCAACATCCTCCAGCTCTTG OTGGCTACTAGATTCTGTAGTTGTTTCCGCATGGGTTTAAAATGTTCAGAAAAATATTTT GTGTTUAAAGTCCTAGGTCAAAAGAAAGTAGTAAATAATGGTACCTGAGCCCCCCATTGT CAAGTGCAAAGTAATTGCACATCGAGTATTTTAACCAAAGCTGCCAGGGTATTCTATCTT GAGCACGATTAGATCTTCAGTCTTGGTTATCAGGATACCGCTGAGGGGCTTGCTGGATCC TANANACAAAAATAGTGTTCATACACATAGGCTACTGACCAGTGTTTTTCGGTGTAAGACG **GTGGACTGCATTTTGATCTTGTATTTTTCTCCTTCAAAGACCTCTGAAAGGTAGATCAGT** AGTAGAGTAAGTGATGATATTGACCGGTAGGTTGAACTTCTTCCAAATAGCGTCAAATAT AGCAGTGAACTTTGATTTCCTCAGAGAAGCCATGGCCAGGAGCTAGGTGGGCAACTGTAT 2033 1440 1740 1500 1380 1320 1260 1560 1980 1920 1860 1800 1680 1620

Figure 2: Predicted amino acid sequence of Mouse partial EDG-5 CDNA. X represents an amino acid which cannot be assigned due to poor sequencing information from direct PCR sequencing

- 1 NTGPVSKTLT VNRWFLRQGL LDTSLTASLA NLLVIAVERH MSIMRMRVHS
- 51 NLTKKRVTLL ILLVWAIAIF WGAVPTLGWN CLCNISACSS LAPIYSRSYL
- 101 IFWTVSNLLA FFIMVAVYVR IYMYVKRKTN VLSPHTSGSI SRRRAPMKLM
- .51 KTVMTVLGAP VVCMTPGLVV LLLDGLNCKQ CNVQHVKXWF LLLALLNSVM
- 01 NPLIYCRSPX PPW

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Figure 3A: Nucleotide sequence of a hEDG-5 cDNA inserted into pcDNA3 (SEQ ID NO:13)

Start and stop codons underlined

1351	1301	1251	1201	1151	1101	1051	1001	951	901	851	801	751	701	159	109	551	501	151	401	351	101	251	201	151	101	51	_
catgcat	cgacgcggcc	AAGCATGGGC	TCCATTAATC	GAATGATTAC	CTGGATGCCT	AGGATAGTAT	CATCCCCTCC	AAGATGATCT	ACCCCATCAT	GCATGTGAAA	CIGGIGGIIC	AGACGGTGAT	GCATACAAGT	TACCTGCGGA	TTTTCTGGAC	CATCTCTGCC	GCCATTTTA	ACCTGACCAA	TATCGCCGTG	CAGGGGCTTC	ACACAGGCCC	AGCTGCTGCC	ATCAAAAACA	TITTCIGCCT	CGATGACTGG	TATGACAAGC	gaattegegg
	gcgaattctt	AGTANAGAGA	ACTGCTAGAT	CIGICICIA	CTYGGCCCAC	TAGCCAAGGT	ACAGTCCTCA	GCTGCTTCTC	CTACTCCTAC	AGGTGGTTCC	TGCCCCTCGA	GACTGTCTTA	GGGTCCATCA	TCTACGTGTA	AGTGTCCAAC	IGCICTICCC	TGGGGGGGGT	AAAGAGGGTG	GAGAGGCACA	TGGACAGTAG	AGTITICAAAA	GATTICTICG	GAAAATTTCA	GITTATTITT	ACAGGAACAA	ACATGGACTT	ccgcgtcgac
	ttgcttttta	GGACCTGCTG	TICTITAAAA	CAMAGCCCAT	CCAGGCCTCC	GCAGTCTGCA	GCAGGAGTGA	TCAGGAGAAC	AAGGACGAGG	TGCTGCTGGC	CGGCCTGAAC	GGGGCGTTTG	GCCGCCGGAG	CGTCAAGAGG	CTCATGGCCT	TGGCCCCCAT	CCCCACACTO	ACACTGCTCA	TGTCAATCAT	CTIGACIGCT	ACTITGACTG	CTGGAATTGC	TITCCCCTTT	TITICTAATT	AGCTTGTGAT	TTTTTATAAT	gttcaCTTCT
	ccctggaaga	CATTTAGAGA	AATTTTTTT	GTACAGTGTT	TCTGGGAAAA	ATAAAAGCAC	CACAGGCAGC	CCAGAGAGGC	ACATGTATGG	GCTGCTCAAC	TGCAGGCAGT	TGGTATGCTG	GACACCCATG	AAAACCAACG	TCCTCATCAT	TTACAGCAGG	GCTGGAATT	TITIGCTIGE	GAGGATGCGG	TCCCTCACCA	TCAACCGCTG	CTATGTATTC	TACTACCTGT	CTCTGGTCAT	TGTTTTGTGT	AGGAGCAACA	CCACAATGAA
	aacacccgag	AAGCACAGGE	CATAGTTTAA	ATTTGAGGTC	GAGCTGTTAA	TTCCTAAACT	CAGTACATAG	OTCCCTCTCG	CACCATGAAG	TCCGTCGTGA	GTGGCGTGCA	GACCCCGGGC	AAGCTAATGA	TCTTGTCTCC	GCTTGTGGTG	AGTTACCTTG	GCCTCTGCAA	CTGGGCCATC	GTCCATAGCA	ACTIGCIGGT	GTTTCTCCGT	CTGATGTTTA	TOGCTAATTT	CGCGGCAGTG	GTTGGGACGT	CTGATACTGT	TGAGTGTCAC

Figure 3B: cDNA sequence of clone pC3-hEDG5#3.4 from the region encoding a hEDG5 polypeptide

105 351 101 251 201 151 155 9 158 108 701 651 109 GTCATCGCGG CAGTGATCAA AAACAGAAAA TTTCATTTCC CCTTCTACTA CAACACTGAT ACTOTCGATO ACTGGACTGG AACAAAGCTT GTGATTGTTT ATGAATGAGT GTCACTATGA CAAGCACATG GACTTTTTTT ATAATAGGAG GAATTGCCTC TGCAACATCT CTGCCTGcTC TTCCCTGGCC CCCATTTACA CACCAACTTG CTGGTTATCG CCGTGGAGAG GCACATGTCA ATCATGAGGA CCTGTTGGCT AATTTGGCTG CTGCCGATTT CTTCGCTGGA ATTGCCTATG CAACGTCTTG TCTCCGCATA CAAGTGGGTC CATCAGCCGC CGGAGGACAC ATCATGGTTG TGGTGTACCT GCGGATCTAC GTGTACGTCA AGAGGAAAAC GCAGGAGTTA CCTTGTTTTC TOGACAGTGT CCAACCTCAT GGCCTTCCTC CTTOTCTOGG CCATCGCCAT TTTTATGGGG GCGGTCCCCA CACTGGGCTG TECCGOTCCA TAGCAACCTO ACCAMANAA GOOTGACACT GCTCATTTTG CGCTGGTTTC TCCGTCAGGG GCTTCTGGAC AGTAGCTTGA CTGCTTCCCT TATTCCTGAT GITTAACACA GGCCCAGITT CAAAAACTIT GACTGTCAAC GCAGCCAGTA CATAGAGGAT AGTATTAGCC AAGGTGCAGT CTGCAATAA TCAACTCCOT COTGAACCCT ATCATCTACT CCTACAAGGA CGAGGACATG TGCTGGACCC CGGGCCTGGT GGTTCTGCTC CTCGACGGCC TGAACTGCAG CCATGAAGCT AATGAAGACG GTAATGACTG TCTTAGGGGGC GTTTGTGGTA GAGGCGTCCC TCTCGCATCC CCTCCACAGT CCTCAGCAGG AGTGACACAG TATGGCACCA TGAAGAAGAT GATCTGCTGC TTCTCTCAGG AGAACCCAG GCAGTGTGGC GTGCAGCATG TGAAAAGGTG GTTCCTGCTG CTGGCGCTGC

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Figure 3C: cDNA sequence of clone pC3-hEDG5#28 from the region encoding a hEDG5 polypeptide.

351 101 201 151 155 501 101 101 IGIGIGIEGO GACGITITIC IGCCIGITIA TITTITITIC TAATICICIG 106 158 108 751 701 51 CAACACTGAT ACTOTCGATG ACTGGACAGG AACAAAGCTT GTGATTGTTT CCTGTTGGCT AATTTAGCTG CTGCCGATTT CTTCGCTGGA ATTGCCTATG GTCATCGCGG CAGTGATCAA AAACAGAAAA TTTCATTTCC CCTTCTACTA TGCGGGTCCA TAGCAACCTG ACCAAAAAGA GGGTGACACT GCTCATTTTG ATGNATGAGT GTCACTATGA CAAGCACATG GACTTTTTTT ATAATAGGGG CACCAACTTG CTGGTTATCG CCGTGGAGAG GCACATGTCA ATCATGAGGA CGCTGGTTTC TCCGTCAGGG GCTTCTGGAC AGTAGCTTGA CTGCTTCCCT TATTCCTGAT GTTTAACACA GGCCCAGTTT CAAAAACTTT GACTGTCAAC GCAGGAGTTA CCTTGTTTTC TGGACAGTGT CCAACCTCAT GGCCTTCCTC GAATTGCCTC TGCAACATCT CTGCCTGCTC TTCCCTGGCC CCCATTTACA CTTGTCTGGG CCATCGCCAT TTTTATGGGG GCGGTCCCCA CACTGGGCLG GCAGCCAGTA CATAGAGGAT AGTATTAGCC AAGGTGCAGT CTGCAATAAA TATGGCACCA TGAAGAAGAT GATCTGCTGC TTCTCTCAGG AGAACCCCAGA TCAACTCCGT CGTGAACCCC ATCATCTACT CCTACAAGGA CGAGGACATG GCAGTGTGGC GTGCAGCATG TGAAAAAGGTG GTTCCTGCTG CTGGCGCTGC TGCTGGACCC CGGGCCTGGT GGTTCTGCTC CTCGACGGCC TGAACTGCAG CCATGAAGCT AATGAAGACG GTGATGACTG TCTTAGGGGC GTTTGTGGTA CAACGTCTTG TCTCCGCALA CAAGTGGGTC CATCAGCCGC CGGAGGACAC ATCATEGITTO TEGTETACCT GEGGATCHAC GTGHACGTCA AGAGGAAAAC GAGGCGTCCC TCTCGCATCC CCTCCACAGT CCTCAGCAGG AGTGACACAG

Figure 4A: Aligned hedg-5 cDNA and predicted amino acid sequence. The first 250 bp of DNA sequence (lower case) is derived from genomic DNA flanking the 5' end of the cDNA insert from clone pC3-hedg-55. Sequences from nt 251-1523 are shown in lower case wherever apparent polymorphisms in different human clones were found. Coding region polymorphisms are detailed in Table 1. One intron exists within the coding region of hedg-5, located between nt 996/997 of the cDNA sequence shown.

181 241 121 101 541 481 23 196 61 ctgtggtaggaggtcaggggctatgtcctggaccaaaggacatttgcactgagacctgac cacaagegaagaagAGGTGTTACTTACTCACAGTGATACTGTTCGTGTACCTGAAAAAAA A RECT TO THE TOTAL OF THE ACT OF gatggcttagtgattttacaaargatatttgtttcttctttaaaatttctttctagg tgaagtccagaagttgagggaactaccctcaatcggtcttgcccgaatctttgtcgttaa acttcaggtcttcaactcccttgatgggagttagccagaacgggcttagaaacagcaatt gacaccatcctccagtccccgatacaggacctggtttcctgtaaacgtgactctggactg grggaaggattggactcgccggatcggaccctttgtttgttaattttacacgcgatttac C V G T F F C L F I F F S N S L V I A A TGTGTGTGGGACGTTTTTCTGCCTGTTATTTTTTTTCTAATTCTCTGGTCATCGGG TATTATCCTCGTTGTGACTATGACAGCTACTGACCTGTCCTTGTTTCGAACACTAACAAA ATAATAGGAGCAACACTGATACTGTCGATGACTGGACAGGAACAAAGCTTGTGATTGTTT K T L T V N R N F L R Q G L L D S S L T CAMANACTTTGACTGTCAACCGCTGGTTTCTCCGTCAGGGGCTTCTGGACAGTAGCTTGA GTCACTAGTTTTTTTTTTTTTAAAGTAAAGGGGAAAATGATGGACAACCGATTAAATCGAC GTTTTTGAAACTGACAGTTGGCGACCAAAGAGGCAGTCCCCGAAGACCTGTCATCGAACT GACGGCTAAAGAAGCGACCTTAACGGATACATAAGGACTACAAATTGTGTCCGGGTCAAA CAGTGATCAAAAACAGAAAATTTCATTTCCCCCTTTTACTACCTGTTGGCTAATTTAGCTG CTGCCGATTTCTTCGCTGGAATTGCCTATGTATTCCTGATGTTTAACACAGGCCCAGTTT VIKNRKPHPPPYYLLANLAA NRSNIDIVDUNICIKLVIVL A D P P A G I A Y V F L H F N T G P V S S S L T . 540 180 120 60 . 600 240 360 90 480 120

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Figure 4A (cont.)

1320	S Q Y I E D S I S Q G A V C N K S T S • GCACCAGTACATAGAGGATAGTATTAGCCAAGGTGCAGTCAGT	1261
1260	N P E R R P S R I P S T V L S R S D T G AGAACCCAGAGAGAGGTCCCCTCCACAGTCCTCAGCAGGAGTGACACAG TCTTGGGTCTCTCCGCAGGGAGAGCGTAGGGGAAGGTGTCAGGAAGTCGTCCTCACTGTGTC	1201
1200	Y K D E D M Y G T M K K H I C C P S Q E CCTACAAGAACAACAACAACAACAACACCATGAAGAAAGA	wi
1140	K R W F L L L A L L N S V V N P I I Y S TGAAAAGGTGGTTCCTGCTGGCGCTGGCGCTCAACTCCGTCGTGAACCCCCATCATCTACT ACTTTTCCACCAAGGACGACGACGACGACGAGTTGAGGCAGCACTTGGGGTAGTAGATGA	1061
1080	G L V V L P L D G L N C R Q C G V Q H V CGGGCCTGGATGGTTCTGCCCCTCGACGGCTGAACTGCAGGCAG	1021
1020	H K L H K T V H T V L G A F V V C H T P CCATGAAGCTAATGAAGACGGTGATGACCGCTACTGACAGAATCCCCGGCAAACACCATACGACCTGGGGGGTACTCGACAAATCCCCGCAAACACCATACGACCTGGG	961
960	R K T N V L S P H T S Q S I S R R R T P AGAGGAAAACCGACCTCTTGTCTCCCCCATACAAGTGGGTCCATCAGCCGCCGCCGAGGACACCTCTTTGTTTCACCCTTTTGGTTGCAGAACAGAGGCGTATGTTCACCCAGGTAGTCGGCGGCCTCCTGTG	901
900	N L H A F L I H V V V Y L R I Y V Y V K CCAACCTCATGGCCTTCCTCATCATGGTTTGTGTGTGTGT	643
8	A C S S L A P I Y S R S Y L V F W T V S CTGCCTGCTCTCCCTGGCCCCATTTACAGCAGGAGTTACCTTGTTTTTCTGGACAGTGT GACGGACGAGAAGGGACCGGGGGTAAATGTCCTCCTCAATGGAACAAAAAAAA	781
780	I A I P N G A V P T L G W N C L C N I S CCATCGCCATTITIATGGGGCGGTCCCCACCACTGGACTTGGAATTGCCTCTGCAACATCT GGTAGCGGTAAAAATACCCCCGCCAGGGGTTGTGACCCGACCTTAACGGAGACGTTGTAGA	721
720	R V H S N L T K K R V T L L I L L V H A TGCGGGTCCATAGCAACCTGACCAAAAAGAGGGTGACACTGCTCATTTTGCTTGTCTGGG ACGCCCAGGTATCGTTGGACTGGTTTTTCTCCCACTGTGACGAGTAAAACGAACAGACCC	661
660	A S L T N L L V I A V E R H M S I M R M CTOCTTCACCAACTTGCTGGTTATCGCCGTGGAGAGGCACATGTCAATCATGAGGA GACGAAGGGAGTGGTTAGAGCGACCAATAGCGGCACCTCTCCGTGTACAGTTAGTACTCCT	601

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Figure 4A (cont.)

1381 1321 1501 1441 AAACTCTGGGATGCCTCTYGGCCCACCCAGGCCTCCTCTGGGAAAAGAGCTGTTAAGAATG ATCTAAAGAAATTTTTTAAAAAAAAAGTATCAAATTTTCGTACCCGTCATTTCTCTCCTGG TTTGAGACCTACGGAGARCCGGGTGGGTCCGGAGAGACCCTTTTCTCGACAATTCTTAC ACGACGTAAATCTCTTTCGTGTC TGCTGCATTTAGAGAAAGCACAG TAGATTTCTTTAAAAAATTTTTTTTCATAGTTTAAAAGCATGGGCAGTAAAGAGAGGACC TAATGGACAGAGATTGTTTCGGGTACATGTCACAATAAACTCCAGAGGTAATTAGTGACG ATTACCTGTCTCTAACAAAGCCCATGTACAGTGTTATTTGAGGTCTCCATTAATCACTGC 1523 1500 1440 1380

clone pC3-hEDG5#3.4 Figure 4B: Predicted amino acid sequence of hEDGS encoded by

101 251 201 151 101 51 YGTMKKMICC FSQENPERRP SRIPSTVLSR SDTGSQYIED SISQGAVCHY LVWAIAIFMG AVPTLGWNCL CNISACSSLA PIYSRSYLVF WTVSNLMAFI VIAAVIOORK PHPPPYYLLA NLAAADPFAG IAYVFLMPNT GPVSKTLTVN CHTPGLVVLL LDGLNCRQCG VQHVKRWFLL LALLNSVVNP IIYSYKDEDM IMVVVYLRIY VYVKRKTNVL SPHTSGSISR RRTPMKLMKT VMTVLGAFVV MNECHYDKHM DFFYNRSNTD TVDDWTGTKL VIVLCVGTPF CLFIFFSNSI SSLTASLINL LVIAVERHMS IMRMRVHSNL TKKRVTLLIL

clone pC3-hEDG5#28. Figure 4C: Predicted amino acid sequence of hEDG5 encoded by

351

STS

101 251 201 151 101 51 VIAAVIOURK PHEPFYYLLA NLAAADPPAG IAYVFLMFNT GPVSKTLTVN MNECHYDIGM DEFYNRONTD TVDDWTGTKL VIVLCVGTFF CLFIFFSNSL CHTPGLVVLL LDGLNCRQCG VQHVKRWFLL LALLNSVVNP IIYSYKDEDN LVWAIAIFMG RWFLRQGLLD SSLTASLTNL LVIAVERHMS IMRMRVHSNL TKKRVTLLIL YGTMKJOHICC FSQENPERRP SRIPSTVLSR SDTGSQYIED SISQGAVCNI VYVKRKTNVL SPHTSGSISR RRTPMKLMKT VMTVLGAFVV AVPTLOWNCL CNISACSSLA PIYSRSYLVF WTVSNLMAFL

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Pigure 5A: Alignment of predicted amino acid sequences of HEDG5 translation products of clones pC3-hEdg5#3.4, pC3-hEdg5. and pC3-hEdg5#28. Amino acid substitutions are indicated in reverse bold text.

he5#3.4_981216 heedg5 he5#28_981215 hsedg5 he5#28_981215 hsedg5 he5#28_981215 he5#3.4_981216 he5#3.4_981216 he5#3.4_981216 hsedg5 he5#3.4_981216 hsedg5 he5#3.4_981216 he5#28_981215 he5#3.4_981216 he5#3.4_981216 he5#28_981215 he5#28_981215 he5#28_981215 haedg5 1
NRECHYDKIM DEFYNRSNTD TVDDWTGTKL VIVLCVGTEF CLEIFFSNSL MNECHYDKIM DEFYNRSNTD TVDDWTGTKL VIVLCVGTEF CLEIFFSNSL MNECHYDKIM DEFYNRSNTD TVDDWTGTKL VIVLCVGTEF CLEIFFSNSL VIAAVIKHRK PHPPPYYLLA NLAAADFPAG IAYVFLMFNT GPVSKTLTVN VIAAVIKHK EHPPPYYLLA NLAAADFPAG IAYVFLMFNT GPVSKTLTVN VIAAVIKHK EHPPPYYLLA NLAAADFPAG IAYVFLMFNT GPVSKTLTVN CMIDGIANTE CMIDGIANTE LVWAIAIFMG AVPTLGWNCL CNISACSSLA PIYSRSYLVF WTVSNLMAFL LVWAIAIFMG AVPTLGWNCL CNISACSSLA PIYSRSYLVF WTVSNLMAFL LVWAIAIFHG AVPTLGWNCL CNISACSSLA PIYSRSYLVF WTVSNLMAFL RWPLRQGLLD SSLTASLTNI LVIAVERHMS IMRMRVHSNI TKKRVTLLIL RWPLRQGLLD SSLTASLTNI LVIAVERHMS IMRMRVHSNI TKKRVTLLIL RWPLRQGLLD SSLTASLTNI LVIAVERHMS IMRMRVHSNI TKKRVTLLIL 351 STS-STS-YGTMKIMICC FSQENPERRP SRIPSTVLSR SDTGSQYIED SISGGAVCHK YGTMKIMICC FSQENPERRP SRIPSTVLSR SDTGSQYIED SISGGAVCHK YGTMKIMICC FSQENPERRP SRIPSTVLSR SDTGSQYIED SISGGAVCHK IMVVVYLRIY VYVKRKTNVL SPHTSGSISR RRTPMKLMKT VMTVLGAFVV IMVVVYLRIY VYVKRKTNVL SPHTSGSISR RRTPMKLMKT VMTVLGAFVV IMVVVYLRIY VYVKRKTNVL SPHTSGSISR RRTPMKLMKT VMTVLGAFVV 151 LDGLNCRQCG VQHVKRWFLL LALLNSVVNP IIYSYKDEDM LDGLNCRQCG VQHVKRWFLL LALLNSVVNP IIYSYKDEDM LDGLNCRQCG VQHVKRWFLL LALLNSVVNP IIYSYKDEDM

he5#28_981215

WO 99/33972 Prigure 5B: Alignment of the amino acid sequence of the murine edg-5 with the amino acid sequence of the human edg-5 of the pC3-hEdg5-3.4 clone.

	SCORES Smith-Wate
	Initl: rman score
10	1981 Init
20	:n: 215 91.2 1
30	5 Opt: identity
6	SCORES Initl: 1981 Initn: 2155 Opt: 2170 z-score: 384.5 E(): 2e-16 Smith-Waterman score: 2170; 91.2% identity in 354 aa overlap
50	e: 384.5 E verlap
60	:(): 2e-16

MEDG5 HEDG5-3.4	MEDG5	MEDGS HEDG5-3.4	MEDG5	NEDG5	MEDCS
310 320 340 350 THINRWICCALODSHTERRPSRAPSTIHSRSETGSOYLEDSISOGPVCNONGS - - -	250 260 270 280 290 300 260 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270 280 270	240 210 220 230 240 25 PIYSRSYLIFMTVSNILLAFFINVAVVRLYMVKRKTRVLSPHTSGSISGRRAPHKLJKT	130 140 150 160 170 180 35 LVIAVERHNSIMRMRVHSNLTKKRVTLLILLVMAIAIFMGAVPTLGMNCLCNISACSSLA	70 80 120 120 120 120 120 120 120 120 120 12	10 50 60 NYBECHYDKHADFFYNRSNYDTADEWTGTLLVIVLCVGTFFCLFIFFSNSLIVIAAVITNRK 11

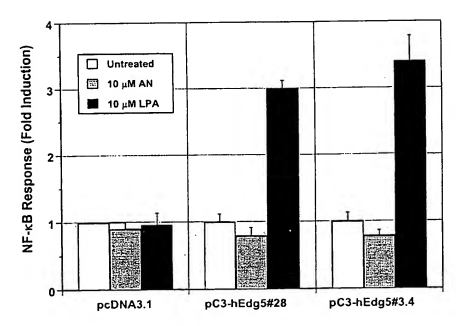


Figure 6

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SEQUENCE LISTINGS

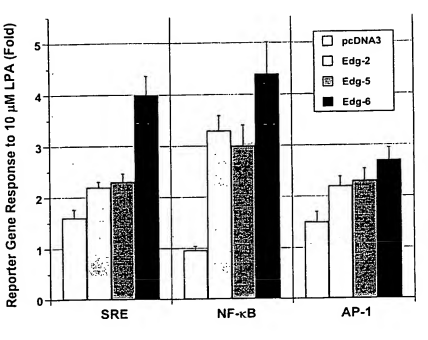


Figure 7 13/13

> (1) GENERAL INFORMATION: (iv) CORRESPONDENCE ADDRESS:
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> (B) STREET: Suite 1500, P.O. Box 190
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>
> Toronto-Dominion Centre (VI) PRIOR APPLICATION DATA:
>
> (A) COUNTRY: U.S.A.
>
> (B) APPLICATION NUMBER: 08/997,803
>
> (C) FILING DATE: 24-DEC-1997
>
> (D) CLASSIFICATION: (1) APPLICANT:
> NAME: ALLELIX BIOPHARMACEUTICALS INC.
> STREET: 6850 Goreway Drive (vi) CURRENT APPLICATION DATA:
> (A) APPLICATION NUMBER: PCT/CA98/01193
> (B) FILING DATE: 24-DEC-1998
> (C) CLASSIFICATION: (v) COMPUTER READABLE FORM: (111) NUMBER OF SEQUENCES: 15 (ii) TITLE OF INVENTION: MANMALIAN EDG-S RECEPTOR HOMOLOGS CITY: Mississauga PROVINCE: Oncario COUNTRY: Canada POSTAL CODE: L4V 1V7 TELEPHONE: (905) 677-0831 PACSIMILE: (905) 677-9595 (C) CITY: Toronto
> (D) PROVINCE: Ontario
> (E) COUNTRY: Canada
> (F) ZIP: M5K 1H6 (A) MEDIUM TYPE: Floppy disk
> (B) COMPUTER: IBM PC compactble
> (C) OPERATING SYSTEM: PC-DOS/MS-DOS
> (D) SOFTWARE: DOS EDITOR

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CTGNYKWTTC ATNAWNMENT ANAYNAYNGG RTT

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

	165 170 175	2
528	OTO GTG TGG ACC CCG GGT CTG GTG CTG CTG CTG GAC GGC CTG S:	
480	GCT CCC ATG AAG CTA ATG AAG ACA GTG ATG ACC GTC TTA GGC GCC TTC Ala Pro Met Lys Leu Het Lys Thr Val Het Thr Val Leu Gly Ala Phe 145 150	
432	ACC AAC GTC TTA TCT CCA CAC ACC AGT GGC TCC ATC AGC CGC CGG AGG Thr Asn Val Leu Ser Pro His Thr Ser Gly Ser Ile Ser Arg Arg Arg 130 135	
38 4	ATC ATG GTG GCG GTA TAC GTA CGC ATC TAC ATG TAT GTT AAA AGG AAA 36 Ile het Val Ala Val Tyr Val Arg Ile Tyr het Tyr Val Lys Arg Lys 115	
316	TTC TGG ACT GTG TCC AAC CTC CTG GCC TTC TTC Phe Trp Thr Val Ser Aen Leu Leu Ala Phe Phe 105	
288	TGC CTC TGC AAC ATC TCG GCC TGC TGT TGT CTG GCT CGC ATT TAC AGT 28 Cys Lau Cys Asn lle Ser Ala Cys Ser Ser Leu Ala Pro lle Tyr Ser 85 90 95	
240	TGG GCC ATC GCC ATC TTC ATG GGG GCC GTC CCC ACN CTG GGA TGG AAT TTP Ala Ile Ala Ile Phe Met Gly Ala Val Pro Thr Leu Gly Trp Ann 65 70 75	
192	CAC AGC AAC TTO ACC AAA AAG CGG GTG ACG CTG CTC ATT CTG CTG GTG 19 His Ser Asn Leu Thr Lys Lys Arg Val Thr Leu Leu Ile Leu Leu Val 50 55	
ī	CTG GTT ATT GCT GTG GAA AGA CAC ATG TCN ATC ATG AGG AGG AGG GTC 14 Leu val Ile Ala Val Glu Arg His Met Ser Ile Met Arg Met Arg Val 35	
96	CGC CAG GGG CTC CTA GAC ACC AGC CTG ACT GCC TCC CTG GCC AAT TTO 9 Arg Gln Gly Leu Leu Asp Thr Ser Leu Thr Ala Ser Leu Ala Asn Leu 25	
å B	AAC ACT GGC CCG GTG TCG AAA ACG TTG ACC GTC AAC CGC TGG TTC CTC Asn Thr Gly Pro Val Ser Lys Thr Leu Thr Val Asn Arg Trp Phe Leu 1 5	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1634	25
	The contract of the contract o	

(ix) FEATURE:

(A) NAME/KEY: Modified Site

(B) LOCATION:16

(D) OTHER INFORMATION: N is Incoine

(ix) FEATURE:

(A) NAME/KEY: Modified Site
(B) LOCATION:13
(D) OTHER INFORMATION: N is Inosine

(ix) FEATURE:

(A) NAME/KEY: Modified Site
(B) LOCATION:4
(D) OTHER INFORMATION: N is Inosine

(11) MOLECULE TYPE: DNA

(ix) FEATURE:
(A) NAME/KEY: Modified Site
(B) LOCKTION:22
(D) OTHER INFORMATION: N is Inosine

(ix) FEATURE:
(A) NAME/KEY: Modified Site
(B) LOCATION:25
(D) OTHER INFORMATION: N is Inceine

(ix) FEATURE:

(A) HAME/KEY: Modified Site
(B) LOCATION:28
(D) OTHER INFORMATION: N is Inosine

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(2) INFORMATION FOR SEQ ID NO:3:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 639 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

AAYTRSATMT STAAYYTGCG TGCGA (1) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 25 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear (x1) SEQUENCE DESCRIPTION: SEQ ID NO:1: (ii) MOLECULE TYPE: cDNA

ຣ

INFORMATION FOR SEQ ID NO:2:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 33 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(2) INFORMATION FOR SEQ ID NO:1:

WO 99/33972

PCT/CA98/01193

(1) INFORMATION FOR SEQ ID NO:6: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDENESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DHA (xi) SEQUENCE DESCRIPTION, SEQ ID NO:6: ATGCGGCTGC ATAGCAACCT GACCAAAAAG 30	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 37 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5: THITTICTAG ACGGTCATCA CTGTCTTCAT TAGCTTC 37	(2) INPORMATION FOR SEQ ID NO:4: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 37 base pairs (8) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4: TITITACTCG AGAITTGCTG GITATTGCTG TGGAAAG 37	TGC Cys	WO 99/33972 PCT/CA98/01193 AAC TGC AAG CAG TGT AAC GTG CAA CAC GTG AAG NGC TGG TTC CTG CTG 576
(2) INFORMATION FOR SEQ ID NO.10: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDENESS: single (D) TOPOLOGY: linear (ii) HOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10: GGGTAGTCGG TACCTCTAGA GCAAGTTCAG CC	(2) INFORMATION FOR SEQ ID NO:9: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic scid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:	(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (11) MOLECULE TYPE: DNA (x1) SEQUENCE DESCRIPTION: SEQ ID NO:8: TITTGAGCAA GTTCAGCCTG GTTAAGT	(1) SEQUENCE CHARACTER SILES: (A) LENGTH: 30 base pair (B) TYPE: nucleic acid (C) STRANDEDEES; single (D) TOPOLOGY: linear (Ai) MOLECULE TYPE: DNA (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: (Xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:	WO 99/33972 (2) THFORMATION FOR SEQ ID NO:7:
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(2) INFORMATION FOR SEQ ID NO:11:

Ξ

(A) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 31 base pairs

(B) TYPE: nucleic acid

(C) STRANDENNESS: single

(D) TOPOLOGY: linear

TIT ATT TIT TOT AAT TOT CTG GTC ATC GCG GCA GTG ATC AAA AAC	GGA ACA ANG CTT GTO ATT GTT TTG TGT GTT GGG ACG TTT TTC TGC CTG Gly Thr Lys Leu Val Ile Val Leu Cys Val Gly Thr Phe Phe Cys Leu 240 245	OAC TIT TAT AAT AGG AGG AAC ACT GAT ACT GTC GAT GAC TOG ACA Aup Phe Phe Tyr Aen Arg Ser Aen Thr Aep Thr Vel Aep Aep Trp Thr 225	ATGITCACIT CTTCTCCACA AIG AAT GAG TOT CAC TAT GAC AAG CAC AIG Met Asn Glu Cys His Tyr Asp Lys His Met 215	OATGGCTIAG TGACTGATTT TACAAATGAT ATTTGTTTCT TCTTTAAATT TCTTTCTAGG	ACTICAGGIC TICAACTCCC TIGAIGGGAG TIAGCCAGAA CGGGCITAGA AACAGCAAIT	CTOTGGTAGG AGGTCAGGGG CTATGTCCTG GACCAAAGGA CATTTGCACT GAGACCTGAC	CACCITICETA ACCIDADEGG CCTAGECTGO GAAACAAACA ATTAAAATGI GEGETAAAIG
434	386	336	290	240	180	120	60

E (× (11 FEATURE MOLECULE TYPE: DNA PEATURE: (A) NAME/KEY: CDS
(B) LOCATION: 261..1322

(2) INFORMATION FOR SEQ ID NO:12:

ATAACAGAGG ATCCTCGAGT ATTTCTTCCA G (xi) SEQUENCE DESCRIPTION: SEQ ID

NO:11:

ב

(11) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1523 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear (A) NAME/KEY: Termination codon
(B) LOCATION: 1320...1322

AAC CTG ACC A Asn Leu Thr L 350 i A TGC TGG Cys Trp ATG AAG GTC TTG Val Leu TAC CTT **3** 100 ATC AGG CAG TOT GGC GTG CAG CAT GTG AAA AGG TGG TTC CTG Arg Gln Cys Gly Val Gln His Val Lys Arg Trp Phe Leu 485 TED TA GCC GTG GAG AAC ATC : val val Ale Ile p Thr Pro Gly Leu 465 Val OO CTA ATG AAG ACG GTG Leu Met Lys Thr Val 450 TCT CCG CAT ACA AGT GGG TCC ATC AGC CGC CGG Ser Pro Hie Thr Ser Gly Ser Ile Ser Arg Arg 435 C TCT GCC TGC TCT 1 e Ser Ala Cys Ser S 385 Lys Lys TIT ATO GGG GCG GTC CCC ACA CTG GGC TGG
Phe Mat Gly Ala Val Pro Thr Leu Gly Trp
370 TAC CTG CGG ATC TAC GTG TAC GTC AAG AGG Tyr Leu Arg Ile Tyr Val Tyr Val Lys Arg 420 Phe Phe AGG TGG ACA 3 AGG GTG 8 Arg Val 355 3 CAC ATG Val Val ¥ 276 r Ser Leu S TCC AAC ACA CTG CTC ATT TTG Thr Leu Leu Ile Leu 360 TCA ATC ATG AGG ATG Ser Ile Met Arg Met 345 ATG ACT GTC TTA GGG GCG Met Thr Val Leu Gly Ala 455 CTG CCC CTC GAC GGC Leu Pro Leu Asp Gly 470 Ala Pro F C Met ATT 11e Ala 7 7 AC Arg CGG 15 C.L.L. Phe 10 1 Leu Leu C CTG AAC Leu Asn 475 agg ACA CCC a Arg Thr Pro AAA ACC AAC Lys Thr Asn CTC ATC AAT TGC CTC ABD CYB Leu 380 Phe Val OTC Ser 395 Trp Ala Arg CAT Val Val 81A 충형 VAI VAI Mec Ber Ber ACC 1106 1154 1058 1010 962 914 866 918 770 722

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Pho lle Pho Pho Ser Asn Ser Leu Val Ile Ala Ala Val Ile Lys Asn 255 260 265

AGA AAA Arg Lys 270 Asp Phe Phe CAT TTC Pho T GGA ATT Pro Phe 7 K A1A TAC CTG TTG GCT AST TTA Met Pho GCT GCT Ala Ala 285

530

482

290 c TAT GTA TTC CTG I a Tyr Val Phe Leu P 295 RAC ACA
ABR Thr

GGC CCA GIT TCA AAA ACT TIG ACT GIC AAC CGC TGG Gly Pro Val Ser Lys Thr Leu Thr Val Asn Arg Trp 305 p Phe Leu Arg Gln
315

578

GGG CTT CTG GAC AGT AGC TTG ACT GCT TCC CTC ACC AAC TTG Gly Leu Leu Asp Ser Ser Leu Thr Ala Ser Leu Thr Asn Leu 320 CTG GTT 626

674

CTG CTC AAC TCC GTG GTG AAC CCC ATC ATC TAC TCC TAC Leu Leu Asn Ser Val Val Asn Pro Ile Ile Tyr Ser Tyr C Lys Asp Glu

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GAC ATG TAT GGC ACC ATG AAG AAG ATG ATG TGC TGC TTC TCT CAG GAG Asp Met Tyr Gly Thr Met Lys Lys Met Ile Cys Cys Phe Ser Gln Glu 510 525 돌 AMC CCA GAG Asp Thr Gly Ser Gln Tyr Ile 545 各점 195 AAT AAA AGC ACT TCC TAA ACTCTGGATG CCTCTYGGCC CACCCAGGCC Asn Lys Ser Thr Ser Arg g CGT CCC TCT CGC ! g Arg Pro Ser Arg 1 530 500 A GAG GAT AGT . e Glu Asp Ser : 550) Ile Pro Ser 535 r ATT AGC CAA r Ile Ser Gln 555 ACA GTC E C GOT GCA Ser Arg 1352 1298 1250 1202

TAAAAGCATG GGCAGTAAAG AGAGGACCTG CTGCATTTAG AGAAAGCACA G GTTATTTGAG GTCTCCATTA ATCACTGCTA GATTTCTTTA AAAAATTTTT TTTCATAGTT TECTETGGGA AAAGAGETGT TAAGAATGAT TACETGTETE TAACAAAGCE CATGTACAGT

> 1472 1412

1523

:2 INFORMATION FOR SEQ ID NO:13

E (A) LENGTH: 1356 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: 11per

CTCTGGTCAT CGCGGCAGTG ATCAAAAACA GAAAATTTCA TTTCCCCTTT TACTACCTGT AGCTTGTGAT TGTTTTGTGT GTTGGGACGT TTTTCTGCCT GTTTATTTTT TTTTCTAATT ACATGGACTT GAATTCGCGG CCGCGTCGAC GTTCACTTCT CCACAATGAA TGAGTGTCAC TATGACAAGC TOTCAATCAT GAGGATGCGG GTCCATAGCA ACCTGACCAA AAAGAGGGTG ACACTGCTCA TGGACAGTAG CTTGACTGCT TCCCTCACCA ACTTGCTGGT TATCGCCGTG GAGAGGCACA TITICIGGAC AGIGICCAAC CICAIGGCCT ICCTCAICAI GGIIGIGGIG TACCIGCGGA GCCTCTGCAA CATCTCTGCC TGCTCTTCCC TGGCCCCCAT TTACAGCAGG AGTTACCTTG TITIGCITGI CIGGGCCAIC GCCATTITTA IGGGGGCGGI CCCCACACIG GGCIGGAAII ACACAGGCCC AGTITICAAAA ACTITIGACTG TCAACCGCTG GTTTCTCCGT CAGGGGCTTC IGGCTAATTT AGCTGCTGCC GATTTCTTCG CTGGAATTGC CTATGTATTC CTGATGTTTA TCTACGTGTA CGTCAAGAGG AAAACCAACG TCTTGTCTCC GCATACAAGT GGGTCCATCA TTTTTATAAT AGGAGCAACA CTGATACTGT CGATGACTGG ACAGGAACAA 540 900 240 120 600 480 120 360

(11) MOLECULE TYPE: DNA

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:13

Ser Leu Val Ile Ala Ala Val Ile Lys Asn Arg Lys Phe His Phe Pro 50 55 60 Val Leu Cys Val Gly Thr Phe Phe Cys Leu Phe Ile Phe Phe Ser Asn 35 40 45 Ser Asn Thx Asp Thr Val Asp Asp Trp Thr Gly Thr Lys Het Asn Glu Cys His Tyr Asp Lys His Met Asp Phe Phe Tyr Asn Arg 1 15 Len Ile Ala Tyr Val Phe Leu Met Phe Asn Thr Gly Pro Val Ser Lys Thr 85 90 95 Leu Leu Phe Tyr Tyr Leu Leu Ala Asn Leu Ala Ala Ala Asp Phe Phe Ala Gly 65 70 80 Met Ser Ile Met Arg Met Arg Val His Scr Asn Leu Thr Lys Lys Arg Thr Ala Ser Leu Thr Asn Leu Leu Val Ile Ala Val Glu Arg His 115 120 126 Thr Val Asn Arg Trp Phe Leu Arg Gln Gly Leu Leu 100 110 1 Leu Val Ile 30

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GTGGCGTGCA GCATGTGAAA AGGTGGTTCC TGCTGCTGGC GCTGCTCAAC TCCGTCGTGA TOGTATOCTO GACCCCOGOC CTOGTGGTTC TGCCCCTCGA CGGCCTGAAC TGCAGGCAGT GCCGCCGGAG GACACCCATG AAGCTAATGA AGACGGTGAT GACTGTCTTA GGGGCGTTTG

GCAGGAGTGA CACAGGCAGC CAGTACATAG AGGATAGTAT TAGCCAAGGT GCAGTCTGCA SCIECTICIC TCAGGAGAAC CCAGAGAGGC GICCCICICG CAICCCCICC ACAGICCICA ACCCCATCAT CTACTCCTAC AAGGACGAGG ACATGTATGG CACCATGAAG AAGATGATCT

1080 1020 960 900

ATAAAAGCAC TICCIAAACT CIGGAIGCCI CIGGCCCACC CAGGCCICCI CIGGGAAAAG

AGCTGTTAAG AATGATTACC TGTCTCTAAC AAAGCCCATG TACAGTGTTA TTTGAGGTCT 1200 1140

GTAAAGAGAG GACCTGCTGC ATTTAGAGAA AGCACAGGTC GACGCGGCCG CGAATTCTTT CCATTAATCA CTGCTAGATT TCTTTAAAAA ATTTTTTTTC ATAGTTTAAA AGCATGGGCA 1320 1260

TGCTTTTTAC CCTGGAAGAA ATACTCGAGC ATGCAT

(i) SEQUENCE CHARACTERISTICS:

(2) INFORMATION FOR SEQ ID NO:14:

(A) LENGTH: 353 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear

(11) MOLECULE TYPE: protein

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:14:

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Arg Gln Gly Leu Leu Asp Thr Ser Leu Thr Ala Ser Leu Ala Asn Leu

Asn Thr Gly Pro Val Ser Lys Thr Leu Thr Val Asn Arg Trp Phe Leu 1 5 10 (x1) SEQUENCE DESCRIPTION: SEQ ID NO:15:

(11) MOLECULE TYPE: protein

SEQUENCE CHARACTERISTICS:

(A) LENGTH: 213 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

2 INFORMATION FOR SEQ ID NO:15:

3

Ser

Val Val Leu Pro Leu Asp Oly Leu Asn Cys Arg Oln Cys Oly Val Oln
260 265 Val Met Thr Val Leu Gly Ala Phe Val Val Cys Trp Thr Pro Gly Leu 255 Ser Arg Ile Pro Ser Thr Val Leu Ser Arg Ser Asp Thr Gly Ser Gln
125
136 Lys Lys Met Ile Cys Cys Phe Ser Gln Glu Asn Pro Glu Arg Arg Pro 105 310 315 Tyr Ile Glu Aap Ser Ile Ser Gln Gly Ala Val Cys Asn Lys Ser Thr 340 350

Asn Pro Ile Ile Tyr Ser Tyr Lys Asp Glu Asp Met Tyr Gly Thr Met 290 295 100 His Val Lys Arg Trp Phe Leu Leu Leu Ala Leu Leu Asn Ser Val Val 275 280 285

Ser Gly Ser Ile Ser Arg Arg Arg Thr Pro Met Lys Leu Met Lys Thr 225 230 230

Ile Met Val Ala Val Tyr Val Arg Ile Tyr Met Tyr Val Lys Arg Lys 115 120 120

Thr Asn Val Leu Ser Pro His Thr Ser Gly Ser Ile Ser Arg Arg Arg 130

Arg Ser Tyr Leu Ile Phe Trp Thr Val Ser Asn Leu Leu Ala Phe Phe 100 105

Val Val Cys Trp Thr Pro Gly Leu Val Val Leu Leu Leu Asp Gly Leu 165

Asn Cys Lys Gln Cys Asn Val Gln His Val Lys Xaa Trp Phe Leu Leu 180 185 190

Pro Xaa Phe Pro Trp 210

Leu Ala Leu Leu Ann Ser Val Met Ann Pro Leu Ile Tyr Cyn Arg Ser 200 205

Ala Pro Met Lys Leu Met Lys Thr Val Met Thr Val Leu Gly Ala Phe 145 150

Cys Leu Cys Asn Ile Ser Ala Cys Ser Ser Leu Ala Pro Ile Tyr Ser 85 90 95

Trp Ala Ile Ala Ile Phe Met Gly Ala Val Pro Thr Leu Gly Trp Asn 65 70 76 80

His Ser Asn Leu Thr Lys Lys Arg Val Thr Leu Leu Ile Leu Leu Val 50 60

Leu Val Ile Ala Val Glu Arg His Met Ser Ile Met Arg Met Arg Val 35 40 45

Ile Tyr Val Tyr Val Lys Arg Lys Thr Asn Val Leu Ser Pro His Thr 210 215

Val Ser Asn Leu Met Ala Phe Leu Ile Met Val Val Val Tyr Leu Arg 195 200 205

Ser Ser Leu Ala Pro Ile Tyr Ser Arg Ser Tyr Leu Val Phe Trp Thr

Ala Val Pro Thr Lou Gly Trp Asn Cys Lou Cys Asn Ile Ser Ala Cys
170
175

Val Thr Leu Leu Ile Leu Leu Val Trp Ala Ile Ala Ile Phe Met Gly 145 150 115 140

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OLIVEIRA L. ET AL.: "A common motif in G-protein-coupled seven transmembrane heltx receptors" JOURNAL OF COMPUTER-AIDED MOLECULAR DESIGN, vol. 7, no. 6, 1 December 1993, pages 649-658, XF002050853	LEE N.H. ET AL.: "Molecular biology of G-protein-coupled receptors" ORUG NEWS AND PERSPECTIVES, vol. 6, no. 7, 1 September 1993, pages 488-497, XP000677175	BURNSTOCK G.: "P2 purinoreceptors: historical perspective and classification" CIBA FOUNDATION SYMPOSIUM, vol. 198, 1996, pages 1-34, XP002102150	FEBS LETTERS, vol. 417, no. 3, 17 December 1997, pages 279-282, xP002102182 see abstract	AN S. ET AL.: "Identification of cDNAs encoding two G protein-coupled receptors for lysosphingolipids"	C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT CAMPONY Citation of document, with indication where appropriate, of the relevant passages
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